(43) International Publication Date 20 December 2001 (20.12.2001)

PCT

(10) International Publication Number WO 01/96381 A2

(51) International Patent Classification7: C07K 14/195

(21) International Application Number: PCT/CA01/00838

(22) International Filing Date: 11 June 2001 (11.06.2001)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

60/211,022

12 June 2000 (12.06.2000) US

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(81) Designated States (national): CA, CO.

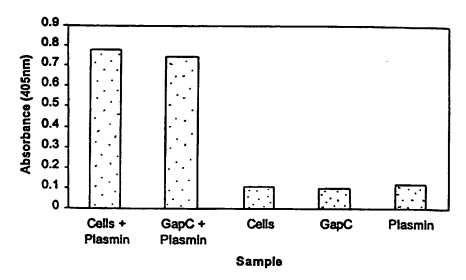
(84) Designated States (regional): European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR).

Published:

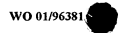
 without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: IMMUNIZATION OF DAIRY CATTLE WITH GapC PROTEIN AGAINST STREPTOCOCCUS INFECTION



(57) Abstract: The GapC plasmin binding protein genes of Streptococcus dysgalactiae (S. dysgalactiae), Streptococcus agalactiae (S. agalactiae). Streptococcus uberis (S. uberis), Streptococcus parauberis (S. parauberis), and Streptococcus iniae (S. iniae) are described, as well as the recombinant production of the GapC proteins therefrom. Also described is the use of the GapC proteins from those species in vaccine compositions to prevent or treat bacterial infections in general, and mastitis in particular.



IMMUNIZATION OF DAIRY CATTLE WITH GapC PROTEIN AGAINST STREPTOCOCCUS INFECTION

TECHNICAL FIELD

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The present invention relates generally to bacterial antigens and genes encoding the same. More particularly, the present invention pertains to the cloning, expression and characterization of the GapC plasmin-binding proteins from Streptococcus dysgalactiae, Streptococcus agalactiae, Streptococcus uberis, Streptococcus parauberis, and Streptococcus iniae, and the use of the same in vaccine compositions.

15 BACKGROUND

Mastitis is an infection of the mammary gland usually caused by bacteria or fungus. The inflammatory response following infection results in decreased milk yield as well as quality, and causes major annual economic losses to the dairy industry.

Among the bacterial species most commonly associated with mastitis are various species of the genus *Streptococcus*, including *Streptococcus aureus*, *Streptococcus uberis* (untypeable), *Streptococcus agalactiae* (Lancefield group B), *Streptococcus dysgalactiae* (Lancefield group C), *Streptococcus zooepidemicus*, and the Lancefield groups D, G, L and N streptococci. Some of those species are contagions (e.g. *S. agalactiae*), while others are considered environmental pathogens (e.g. *S. dysgalactiae* and *S. uberis*).

The environmental pathogen *S. uberis* is responsible for about 20% of all clinical cases of mastitis (Bramley, A.J. and Dodd, F.H. (1984) *J. Dairy Res.* 51:481-512; Bramley, A.J. (1987) *Animal Health Nutrition* 42:12-16; Watts, J.L. (1988) *J. Dairy Sci.* 71:1616-1624); it is the predominant organism isolated from mammary glands during the non-lactating period (Bramley, A.J. (1984) *Br. Vet. J.* 140:328-335; Bramley and Dodd (1984) *J. Dairy Res.* 51:481-512; Oliver, S.P. (1988) *Am. J. Vet. Res.* 49:1789-1793).

Mastitis resulting from infection with *S. uberis* is commonly subclinical, characterized by apparently normal milk with an increase in somatic cell counts due to the influx of leukocytes. The chemical composition of milk is changed due to suppression of

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secretion with the transfer of sodium chloride and bicarbonate from blood to milk, causing a shift of pH to a more alkaline level. *S. uberis* mastitis may also take the form of an acute clinical condition, with obvious signs of disease such as clots or discoloration of the milk and swelling or hardness of the mammary gland. Some cases of the clinical disease can be severe and pyrexia may be present. For a review of the clinical manifestations of *S. uberis* mastitis, see, Bramley (1991) Mastitis: physiology or pathology. p. 3-9. In C. Burvenich, G. Vandeputte-van Messom, and A. W. Hill (ed.), *New insights into the pathogenesis of mastitis*. Rijksuniversiteit Gent, Belgium; and Schalm et al. (1971) The mastitis complex-A brief summary. p. 1-3. In *Bovine Mastitis*. Lea & Febiger, Philadelphia

Conventional antibacterial control methods such as teat dipping and antibiotic therapy are effective in the control of many types of contagious mastitis, but the environmental organisms typically found in all dairy barns are often resistant to such measures. Vaccination is therefore an attractive strategy to prevent infections of the mammary glands, and has been shown to be beneficial in the case of some contagious mastitis pathogens.

The literature is limited regarding vaccination studies with *S. dysgalactiae* and *S. uberis*, and variable results have been observed. In some cases, immunization has resulted in increased sensitivity to the specific organism and in other cases strain-specific protection has been obtained.

For example, previous studies have shown that primary infection with *S. uberis* can considerably reduce the rate of infection following a second challenge with the same strain (Hill, A.W. (1988) *Res. Vet. Sci.* 44:386-387). Local vaccination with killed *S. uberis* protects the bovine mammary gland against intramammary challenge with the homologous strain (Finch et al. (1994) *Infect. Immun.* 62:3599-3603). Similarly, subcutaneous vaccination with live *S. uberis* has been shown to cause a dramatic modification of the pathogenesis of mastitis with the same strain (Hill et al. (1994) *FEMS Immunol. Med. Microbiol.* 8:109-118). Animals vaccinated in this way shed fewer bacteria in their milk and many quarters remain free of infection.

Nonetheless, vaccination with live or attenuated bacteria can pose risks to the recipient. Further, it is clear that conventional killed vaccines are in general largely ineffective against S. uberis and S. agalactiae, either due to lack of protective antigens on in vitro-grown cells or masking of these antigens by molecular mimicry.

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ting mastitis vaccines against S. agalactiae

The current lack of existing mastitis vaccines against *S. agalactiae* or the contagious streptococcus strains is due at least in part to a lack of knowledge regarding the virulence determinants and protective antigens produced by those organisms which are involved in invasion and protection of the mammary gland (Collins et al. (1988) *J. Dairy Res.* 55:25-32; Leigh et al. (1990) *Res. Vet. Sci.* 49: 85-87; Marshall et al. (1986) *J. Dairy Res.* 53: 507-514).

S. dysgalactiae is known to bind several extracellular and plasma-derived proteins such as fibronectin, fibrinogen, collagen, alpha-II-macroglobulin, IgG, albumin and other compounds. The organism also produces hyaluronidase and fibrinolysin and is capable of adhering to and invading bovine mammary epithelial cells. However, the exact roles of the bacterial components responsible for these phenotypes in pathogenesis is not known.

Similarly, the pathogenesis of *S. uberis* infection is poorly understood. Furthermore, the influence of *S. uberis* virulence factors on host defense mechanisms and mammary gland physiology is not well defined. Known virulence factors associated with *S. uberis* include a hyaluronic acid capsule (Hill, A.W. (1988) *Res. Vet. Sci.* 45:400-404), hyaluronidase (Schaufuss et al. (1989) *Zentralbl. Bakteriol. Ser. A* 271:46-53), R-like protein (Groschup, M.H. and Timoney, J.F. (1993) *Res. Vet. Sci.* 54:124-126), and a cohemolysin, the CAMP factor, also known as UBERIS factor (Skalka, B. and Smola, J. (1981) *Zentralbl. Bakteriol. Ser. A* 249:190-194), R-like protein, plasminogen activator and CAMP factor. However, very little is known of their roles in pathogenicity.

The use of virulence determinants from *Streptococcus* as immunogenic agents has been proposed. For example, the CAMP factor of *S. uberis* has been shown to protect vertebrate subjects from infection by that organism (Jiang, U.S. Patent No. 5,863,543).

The γ antigen of the group B Streptococci strain A909 (ATCC No. 27591) is a component of the c protein marker complex, which additionally comprises an α and β subunit (Boyle, U.S. Patent No. 5,721,339). Subsets of serotype Ia, II, and virtually all serotype Ib cells of group B streptococci, have been reported to express components of the c protein. Use of the γ subunit as an immunogenic agent against infections by Lancefield Group B Streptococcus infection has been proposed. However, its use to prevent or treat bacterial infections in animals, including mastitis in cattle, has not been studied.

The group A streptococcal M protein is considered to be one of the major virulence factors of this organism by virtue of its ability to impede attack by human phagocytes (Lancefield, R.C. (1962) *J. Immunol.* 89:307-313). The bacteria persist in the infected

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tissue until antibodies are produced against the M molecule. Type-specific antibodies to the M protein are able to reverse the antiphagocytic effect of the molecule and allow efficient clearance of the invading organism.

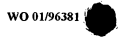
M proteins are one of the key virulence factors of Streptococcus pyogenes, due to their involvement in mediating resistance to phagocytosis (Kehoe, M.A. (1991) Vaccine 9:797-806) and their ability to induce potentially harmful host immune responses via their superantigenicity and their capacity to induce host-cross-reactive antibody responses (Bisno, A.L. (1991) New Engl. J. Med. 325:783-793; Froude et al. (1989) Curr. Top. Microbiol. Immunol. 145:5-26; Stollerman, G.H. (1991) Clin. Immunol. Immunopathol. 61:131-142).

However, obstacles exist to using intact M proteins as vaccines. The protein's opsonic epitopes are extremely type-specific, resulting in narrow, type-specific protection. Further, some M proteins appear to contain epitopes that cross react with tissues of the immunized subject, causing a harmful autoimmune response (See e.g., Dale, J.L. and Beached, G.H. (1982) *J. Exp. Med* 156:1165-1176; Dale, J.L. and Beached, G.H. (1985) *J. Exp. Med*. 161:113-122; Baird, R.W., Bronze, M.S., Drabs, W., Hill, H.R., Veasey, L.G. and Dale, J.L. (1991) *J. Immun*. 146:3132-3137; Bronze, M.S. and Dale, J.L. (1993) *J. Immun* 151:2820-2828; Cunningham, M.W. and Russell, S.M. (1983) *Infect. Immun*. 42:531-538).

Chimeric proteins containing three different fibronectin binding domains (FNBDs) derived from fibronectin binding proteins of *S. dysgalactiae* and *Staphylococcus aureus* have been expressed on the surface of *Staph. carnosus* cells. In the case of one of these proteins, intranasal immunizations with live recombinant *Staph. carnosus* cells expressing the chimeric protein on their surface resulted in an improved antibody response to a model immunogen present within the chimeric surface protein.

A GapC plasmin binding protein from a strain of Group A *Streptococcus* has previously been identified and characterized, and its use in thrombolytic therapies has been described (Boyle, et al., U.S. Patent No. 5,237,050; Boyle, et al., U.S. Patent No. 5,328,996).

However, until now, the protective capability of GapC has not been studied, nor have the GapC proteins of *Streptococcus dysgalactiae*, *Streptococcus agalactiae*, *Streptococcus uberis*, *Streptococcus parauberis* or *Streptococcus iniae* been isolated or characterized.



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Figures 6A-6E shows a DNA alignment created by PileUp and displayed by Pretty software (a component of the GCG Wisconsin Package, version 10, provided by the SeqWeb sequence analysis package, version 1.1, of the Canadian Bioinformatics Resource). The figure depicts the isolated nucleotide sequences of the gapC genes from S. dysgalactiae (DysGapC, Check 9344), S. agalactiae (AgalGapC. Check 2895), S. uberis (UberGapC, Check 5966), S. parauberis (PUberGapC, Check 9672), and S. iniae (IniaeGapC, Check 990). The previously known sequences of S. equisimilis (SeqGapC, Check 5841), S. pyogenes (SpyGapC, Check 4037), and a bovine GAPDH protein (BovGapC, check 5059) are also included. The length and weight parameters were the same for all sequences (1018) and 1.00, respectively). The parameters used in the DNA sequence comparison were as follows: Plurality--2.00; Threshold--1; AveWeight--1.00; AveMatch--1.00; AvMisMatch--0.00; Symbol comparison table--pileupdna.cmp; CompCheck-6876; GapWeight--5; GapLengthWeight--1; PileUp MSF--1018; Type--N; Check--3804. In the figure, dashes represent identical nucleotides; dots represent gaps introduced by the software used to generate the alignment chart, and tildes represent regions not included in the overall alignment due to differences in the length of the gene sequences.

Figures 7A-7B shows an amino acid sequence alignment created by PileUp and displayed by Pretty (as above) that depicts the deduced amino acid sequences of the GapC proteins from S. dysgalactiae (DysGapC, Check 6731), S. agalactiae (AgalGapC, Check 1229), S. uberis (UberGapC, Check 8229), S. parauberis (PUberGapC, Check 8889) and S. iniae (IniaeGapC, check 8785). The previously known sequences of S. equisimilis (SeqGapC, Check 8252), S. pyogenes (SpyGapC, Check 6626) and a bovine GAPDH protein (BovGapC, Check 8479) are also included. In the figure, dashes represent identical amino acid residues; dots represent gaps introduced by the PileUp software, and tildes represent regions not included in the overall alignment due to differences in the length of the gene sequences.

Figure 8 shows Kyte-Doolittle hydropathy plots (averaged over a window of 7), Emini surface probability plots, Karplus-Schulz chain flexibility plots, Jameson-Wolf antigenic index plots, and both Chou-Fasman and Garnier-Osguthorpe-Robson secondary structure plots for the GapC protein isolated from *S. dysgalactiae*.

Figure 9 shows Kyte-Doolittle hydropathy plots (averaged over a window of 7), Emini surface probability plots, Karplus-Schulz chain flexibility plots, Jameson-Wolf



Δ-) represent challenged, Mig-vaccinated animals, and x's (-X-) represent challenged,
 GapC-vaccinated animals.

Figure 26 illustrates somatic cell counts per mammary quarter on day 1 post-challenge. In the figure, the bar represents the mean for each group. Squares (-1) represent unvaccinated animals; triangles (-1) represent GapC-vaccinated animals, and inverted triangles (-1) represent Mig-vaccinated animals.

Figure 27 depicts the somatic cell counts of non-vaccinated, non-challenged and GapC-immunized, high-titer animals (i.e., the four animals exhibiting the highest antibody titres of the eight animals in the particular group) for seven days post challenge, plotted as the log₁₀ of the mean somatic cell count/ml milk against time in days post challenge. Diamonds (-♦-) represent unchallenged, unvaccinated animals, squares (-■-) represent challenged, unvaccinated animals, and triangles (-△-) represent challenged, GapC-vaccinated animals.

DETAILED DESCRIPTION OF THE INVENTION

The practice of the present invention will employ, unless otherwise indicated, conventional techniques of molecular biology, microbiology, recombinant DNA technology, and immunology, which are within the skill of the art. Such techniques are explained fully in the literature. See, e.g., Sambrook, Fritsch & Maniatis, Molecular Cloning: A Laboratory Manual, Vols. I, II and III, Second Edition (1989); Perbal, B., A Practical Guide to Molecular Cloning (1984); the series, Methods In Enzymology (S. Colowick and N. Kaplan eds., Academic Press, Inc.); and Handbook of Experimental Immunology, Vols. I-IV (D.M. Weir and C.C. Blackwell eds., 1986, Blackwell Scientific Publications).

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The following amino acid abbreviations are used throughout the text:

Alanine: Ala (A)	Arginine: Arg (R)
Asparagine: Asn (N)	Aspartic acid: Asp (D)
Cysteine: Cys (C)	Glutamine: Gln (Q)
Glutamic acid: Glu (E)	Glycine: Gly (G)
Histidine: His (H)	Isoleucine: Ile (I)
Leucine: Leu (L)	Lysine: Lys (K)
Methionine: Met (M)	Phenylalanine: Phe (F)

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"Wild type" or "native" proteins or polypeptides refer to proteins or polypeptides isolated from the source in which the proteins naturally occur. "Recombinant" polypeptides refer to polypeptides produced by recombinant DNA techniques; i.e., produced from cells transformed by an exogenous DNA construct encoding the desired polypeptide. "Synthetic" polypeptides are those prepared by chemical synthesis.

An "isolated" protein or polypeptide is a protein or polypeptide molecule separate and discrete from the whole organism with which the molecule is found in nature; or a protein or polypeptide devoid, in whole or part, of sequences normally associated with it in nature; or a sequence, as it exists in nature, but having heterologous sequences (as defined below) in association therewith.

The term "functionally equivalent" intends that the amino acid sequence of a GapC plasmin-binding protein is one that will elicit a substantially equivalent or enhanced immunological response, as defined above, as compared to the response elicited by a GapC plasmin-binding protein having identity with the reference GapC plasmin-binding protein, or an immunogenic portion thereof.

The term "epitope" refers to the site on an antigen or hapten to which specific B cells and/or T cells respond. The term is also used interchangeably with "antigenic determinant" or "antigenic determinant site." Antibodies that recognize the same epitope can be identified in a simple immunoassay showing the ability of one antibody to block the binding of another antibody to a target antigen.

The terms "immunogenic" protein or polypeptide refer to an amino acid sequence which elicits an immunological response as described above. An "immunogenic" protein or polypeptide, as used herein, includes the full-length sequence of the GapC plasmin-binding protein in question, with or without the signal sequence, membrane anchor domain and/or plasmin-binding domain, analogs thereof, or immunogenic fragments thereof. By "immunogenic fragment" is meant a fragment of a GapC plasmin-binding protein which includes one or more epitopes and thus elicits the immunological response described above. Such fragments can be identified using any number of epitope mapping techniques, well known in the art. See, e.g., *Epitope Mapping Protocols* in Methods in Molecular Biology, Vol. 66 (Glenn E. Morris, Ed., 1996) Humana Press, Totowa, New Jersey. For example, linear epitopes may be determined by e.g., concurrently synthesizing large numbers of peptides on solid supports, the peptides corresponding to portions of the protein molecule, and reacting the peptides with antibodies while the peptides are still attached to the

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supports. Such techniques are known in the art and described in, e.g., U.S. Patent No. 4,708,871; Geysen et al. (1984) *Proc. Natl. Acad. Sci. USA* 81:3998-4002; Geysen et al. (1986) *Molec. Immunol.* 23:709-715. Similarly, conformational epitopes are readily identified by determining spatial conformation of amino acids such as by, e.g., x-ray crystallography and 2-dimensional nuclear magnetic resonance. See, e.g., *Epitope Mapping Protocols*, *supra*. Antigenic regions of proteins can also be identified using standard antigenicity and hydropathy plots, such as those calculated using, e.g., the Omiga version 1.0 software program available from the Oxford Molecular Group. This computer program employs the Hopp/Woods method, Hopp et al., *Proc. Natl. Acad. Sci USA* (1981) 78:3824-3828 for determining antigenicity profiles, and the Kyte-Doolittle technique, Kyte et al., *J. Mol. Biol.* (1982) 157:105-132 for hydropathy plots. Figures 8 to 12 herein depict Kyte-Doolittle profiles for representative proteins encompassed by the invention.

Immunogenic fragments, for purposes of the present invention, will usually include at least about 3 amino acids, preferably at least about 5 amino acids, more preferably at least about 10-15 amino acids, and most preferably 25 or more amino acids, of the parent GapC plasmin-binding-binding protein molecule. There is no critical upper limit to the length of the fragment, which may comprise nearly the full-length of the protein sequence, or even a fusion protein comprising two or more epitopes of GapC.

An "immunogenic composition" is a composition that comprises an antigenic molecule where administration of the composition to a subject results in the development in the subject of a humoral and/or a cellular immune response to the antigenic molecule of interest.

By "subunit vaccine composition" is meant a composition containing at least one immunogenic polypeptide, but not all antigens, derived from or homologous to an antigen from a pathogen of interest. Such a composition is substantially free of intact pathogen cells or particles, or the lysate of such cells or particles. Thus, a "subunit vaccine composition" is prepared from at least partially purified (preferably substantially purified) immunogenic polypeptides from the pathogen, or recombinant analogs thereof. A subunit vaccine composition can comprise the subunit antigen or antigens of interest substantially free of other antigens or polypeptides from the pathogen.

By "pharmaceutically acceptable" or "pharmacologically acceptable" is meant a material which is not biologically or otherwise undesirable, i.e., the material may be administered to an individual in a formulation or composition without causing any

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undesirable biological effects or interacting in a deleterious manner with any of the components of the composition in which it is contained.

An "immunological response" to a composition or vaccine is the development in the host of a cellular and/ or antibody-mediated immune response to the composition or vaccine of interest. Usually, an "immunological response" includes but is not limited to one or more of the following effects: the production of antibodies, B cells, helper T cells, suppressor T cells, and/or cytotoxic T cells and/or $\gamma\delta$ T cells, directed specifically to an antigen or antigens included in the composition or vaccine of interest. Preferably, the host will display either a therapeutic or protective immunological response such that resistance of the mammary gland to new infection will be enhanced and/or the clinical severity of the disease reduced. Such protection will be demonstrated by either a reduction or lack of symptoms normally displayed by an infected host and/or a quicker recovery time.

By "nucleic acid immunization" is meant the introduction of a nucleic acid molecule encoding one or more selected antigens into a host cell, for the *in vivo* expression of an antigen, antigens, an epitope, or epitopes. The nucleic acid molecule can be introduced directly into a recipient subject, such as by injection, inhalation, oral, intranasal and mucosal administration, or the like, or can be introduced *ex vivo*, into cells which have been removed from the host. In the latter case, the transformed cells are reintroduced into the subject where an immune response can be mounted against the antigen encoded by the nucleic acid molecule.

The term "treatment" as used herein refers to either (1) the prevention of infection or reinfection (prophylaxis), or (2) the reduction or elimination of symptoms of the disease of interest (therapy).

By "mastitis" is meant an inflammation of the mammary gland in mammals, including in cows, ewes, goats, sows, mares, and the like, caused by the presence of S. uberis. The infection manifests itself by the infiltration of phagocytic cells in the gland. Generally, 4 clinical types of mastitis are recognized: (1) peracute, associated with swelling, heat, pain, and abnormal secretion in the gland and accompanied by fever and other signs of systemic disturbance, such as marked depression, rapid weak pulse, sunken eyes, weakness and complete anorexia; (2) acute, with changes in the gland similar to those above but where fever, anorexia and depression are slight to moderate; (3) subacute, where no systemic changes are displayed and the changes in the gland and its secretion are less

marked: and (4) subclinical, where the inflammatory reaction is detectable only by standard tests for mastitis.

Standard tests for the detection of mastitis include but are not limited to, the California Mastitis Test, the Wisconsin Mastitis Test, the Nagase test, the electronic cell count and somatic cell counts used to detect a persistently high white blood cell content in milk. In general, a somatic cell count of about 300,000 to about 500,000 cells per ml or higher, in milk will indicate the presence of infection. Thus, a vaccine is considered effective in the treatment and/or prevention of mastitis when, for example, the somatic cell count in milk is retained below about 500,000 cells per ml. For a discussion of mastitis and the diagnosis thereof, see, e.g., *The Merck Veterinary Manual: A Handbook of Diagnosis, Therapy, and Disease Prevention and Control for the Veterinarian*, Merck and Co., Rahway, New Jersey, 1991.

By the terms "vertebrate," "subject," and "vertebrate subject" are meant any member of the subphylum Chordata, including, without limitation, mammals such as cattle, sheep, pigs, goats, horses, and humans; domestic animals such as dogs and cats; and birds, including domestic, wild and game birds such as cocks and hens including chickens, turkeys and other gallinaceous birds; and fish. The term does not denote a particular age. Thus, both adult and newborn animals, as well as fetuses, are intended to be covered.

A "nucleic acid" molecule can include, but is not limited to, procaryotic sequences, eucaryotic mRNA, cDNA from eucaryotic mRNA, genomic DNA sequences from eucaryotic (e.g., mammalian) DNA, and even synthetic DNA sequences. The term also captures sequences that include any of the known base analogs of DNA and RNA.

An "isolated" nucleic acid molecule is a nucleic acid molecule separate and discrete from the whole organism with which the molecule is found in nature; or a nucleic acid molecule devoid, in whole or part, of sequences normally associated with it in nature; or a sequence, as it exists in nature, but having heterologous sequences (as defined below) in association therewith. The term "isolated" in the context of a polynucleotide intends that the polynucleotide is isolated from the chromosome with which it is normally associated, and is isolated from the complete genomic sequence in which it normally occurs.

"Purified polynucleotide" refers to a polynucleotide of interest or fragment thereof which is essentially free, e.g., contains less than about 50%, preferably less than about 70%, and more preferably less than about 90%, of the protein with which the polynucleotide is naturally associated. Techniques for purifying polynucleotides of interest are well-known

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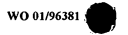


in the art and include, for example, disruption of the cell containing the polynucleotide with a chaotropic agent and separation of the polynucleotide(s) and proteins by ion-exchange chromatography, affinity chromatography and sedimentation according to density.

A "coding sequence" or a "nucleotide sequence encoding" a particular protein, is a nucleotide sequence which is transcribed and translated into a polypeptide *in vitro* or *in vivo* when placed under the control of appropriate regulatory elements. The boundaries of the coding sequence are determined by a start codon at the 5' (amino) terminus and a translation stop codon at the 3' (carboxy) terminus. A coding sequence can include, but is not limited to, procaryotic sequences, cDNA from eucaryotic mRNA, genomic DNA sequences from eucaryotic (e.g., mammalian) DNA, and even synthetic DNA sequences. A transcription termination sequence will usually be located 3' to the coding sequence. A "complementary" sequence is one in which the nitrogenous base at a given nucleotide position is the complement of the nitrogenous base appearing at the same position in the reference sequence. To illustrate, the complement of adenosine is tyrosine, and vice versa; similarly, cytosine is complementary to guanine, and vice versa; hence, the complement of the reference sequence 5'-ATGCTGA-3' would be 5'-TACGACT-3'.

A "wild-type" or "native" sequence, as used herein, refers to polypeptide encoding sequences that are essentially as they are found in nature, e.g., the *S. dysgalactiae* GapC protein encoding sequences depicted in Figures 1A-1B (SEQ ID NO:4).

"Recombinant" as used herein to describe a nucleic acid molecule means a polynucleotide of genomic, cDNA, semisynthetic, or synthetic origin which, by virtue of its origin or manipulation: (1) is not associated with all or a portion of the polynucleotide with which it is associated in nature; and/or (2) is linked to a polynucleotide other than that to which it is linked in nature. The term "recombinant" as used with respect to a protein or polypeptide means a polypeptide produced by expression of a recombinant polynucleotide. "Recombinant host cells," "host cells," "cells," "cell lines," "cell cultures," and other such terms denoting procaryotic microorganisms or eucaryotic cell lines cultured as unicellular entities, are used interchangeably, and refer to cells which can be, or have been, used as recipients for recombinant vectors or other transfer DNA, and include the progeny of the original cell which has been transfected. It is understood that the progeny of a single parental cell may not necessarily be completely identical in morphology or in genomic or total DNA complement to the original parent, due to accidental or deliberate mutation. Progeny of the parental cell which are sufficiently similar to the parent to be characterized



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by the relevant property, such as the presence of a nucleotide sequence encoding a desired peptide, are included in the progeny intended by this definition, and are covered by the above terms.

"Homology" refers to the percent identity between two polynucleotide or two polypeptide moieties. Two DNA, or two polypeptide sequences are "substantially homologous" to each other when the sequences exhibit at least about 80%-85%, preferably at least about 90%, and most preferably at least about 95%-98% sequence identity over a defined length of the molecules. As used herein, substantially homologous also refers to sequences showing complete identity to the specified DNA or polypeptide sequence.

In general, "identity" refers to an exact nucleotide-to-nucleotide or amino acid-toamino acid correspondence of two polynucleotides or polypeptide sequences, respectively. Percent identity can be determined by a direct comparison of the sequence information between two molecules by aligning the sequences, counting the exact number of matches between the two aligned sequences, dividing by the length of the shorter sequence, and multiplying the result by 100. Readily available computer programs can be used to aid in the analysis, such as ALIGN, Dayhoff, M.O. in Atlas of Protein Sequence and Structure M.O. Dayhoff ed., 5 Suppl. 3:353-358, National biomedical Research Foundation. Washington, DC, which adapts the local homology algorithm of Smith and Waterman (1981) Advances in Appl. Math. 2:482-489 for peptide analysis. Programs for determining nucleotide sequence identity are available in the Wisconsin Sequence Analysis Package, Version 8 (available from Genetics Computer Group, Madison, WI) for example, the BESTFIT, FASTA and GAP programs, which also rely on the Smith and Waterman algorithm. These programs are readily utilized with the default parameters recommended by the manufacturer and described in the Wisconsin Sequence Analysis Package referred to above. For example, percent identity of a particular nucleotide sequence to a reference sequence can be determined using the homology algorithm of Smith and Waterman with a default scoring table and a gap penalty of six nucleotide positions.

Another method of establishing percent identity in the context of the present invention is to use the MPSRCH package of programs copyrighted by the University of Edinburgh, developed by John F. Collins and Shane S. Sturrok, and distributed by IntelliGenetics, Inc. (Mountain View, CA). From this suite of packages the Smith-Waterman algorithm can be employed where default parameters are used for the scoring table (for example, gap open penalty of 12, gap extension penalty of one, and a gap of six).

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From the data generated the "Match" value reflects "sequence identity." Other suitable programs for calculating the percent identity or similarity between sequences are generally known in the art, for example, another alignment program is BLAST, used with default parameters. For example, BLASTN and BLASTP can be used using the following default parameters: genetic code = standard; filter = none; strand = both; cutoff = 60; expect = 10; Matrix = BLOSUM62; Descriptions = 50 sequences; sort by = HIGH SCORE; Databases = non-redundant, GenBank + EMBL + DDBJ + PDB + GenBank CDS translations + Swiss protein + Spupdate + PIR. Details of these programs can be found at the following internet address: http://www.ncbi.nlm.gov/cgi-bin/BLAST.

Alternatively, homology can be determined by hybridization of polynucleotides under conditions which form stable duplexes between homologous regions, followed by digestion with single-stranded-specific nuclease(s), and size determination of the digested fragments. DNA sequences that are substantially homologous can be identified in a Southern hybridization experiment under, for example, stringent conditions, as defined for that particular system. Defining appropriate hybridization conditions is within the skill of the art. See, e.g., Sambrook et al., *supra*; *DNA Cloning*, *supra*; *Nucleic Acid Hybridization*, *supra*.

By the term "degenerate variant" is intended a polynucleotide containing changes in the nucleic acid sequence thereof, that encodes a polypeptide having the same amino acid sequence as the polypeptide encoded by the polynucleotide from which the degenerate variant is derived.

Techniques for determining amino acid sequence "similarity" are well known in the art. In general, "similarity" means the exact amino acid to amino acid comparison of two or more polypeptides at the appropriate place, where amino acids are identical or possess similar chemical and/or physical properties such as charge or hydrophobicity. A so-termed "percent similarity" then can be determined between the compared polypeptide sequences. Techniques for determining nucleic acid and amino acid sequence identity also are well known in the art and include determining the nucleotide sequence of the mRNA for that gene (usually via a cDNA intermediate) and determining the amino acid sequence encoded thereby, and comparing this to a second amino acid sequence. In general, "identity" refers to an exact nucleotide to nucleotide or amino acid to amino acid correspondence of two polypucleotides or polypeptide sequences, respectively.

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A "heterologous" region of a DNA construct is an identifiable segment of DNA within or attached to another DNA molecule that is not found in association with the other molecule in nature. Thus, when the heterologous region encodes a bacterial gene, the gene will usually be flanked by DNA that does not flank the bacterial gene in the genome of the source bacteria. Another example of the heterologous coding sequence is a construct where the coding sequence itself is not found in nature (e.g., synthetic sequences having codons different from the native gene). Allelic variation or naturally occurring mutational events do not give rise to a heterologous region of DNA, as used herein.

A "vector" is a replicon, such as a plasmid, phage, or cosmid, to which another DNA segment may be attached so as to bring about the replication of the attached segment. A vector is capable of transferring gene sequences to target cells (e.g., bacterial plasmid vectors, viral vectors, non-viral vectors, particulate carriers, and liposomes).

Typically, the terms "vector construct," "expression vector," "gene expression vector," "gene delivery vector," "gene transfer vector," and "expression cassette" all refer to an assembly which is capable of directing the expression of a sequence or gene of interest. Thus, the terms include cloning and expression vehicles, as well as viral vectors.

These assemblies include a promoter which is operably linked to the sequences or gene(s) of interest. Other control elements may be present as well. The expression cassettes described herein may be contained within a plasmid construct. In addition to the components of the expression cassette, the plasmid construct may also include a bacterial origin of replication, one or more selectable markers, a signal which allows the plasmid construct to exist as single-stranded DNA (e.g., a M13 origin of replication), a multiple cloning site, and a "mammalian" origin of replication (e.g., a SV40 or adenovirus origin of replication).

DNA "control elements" refers collectively to transcription promoters, transcription enhancer elements, transcription termination sequences, polyadenylation sequences (located 3' to the translation stop codon), sequences for optimization of initiation of translation (located 5' to the coding sequence), translation termination sequences, upstream regulatory domains, ribosome binding sites and the like, which collectively provide for the transcription and translation of a coding sequence in a host cell. See e.g., McCaughan et al. (1995) PNAS USA 92:5431-5435; Kochetov et al (1998) FEBS Letts. 440:351-355. Not all of these control sequences need always be present in a recombinant vector so long as the desired gene is capable of being transcribed and translated.

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"Operably linked" refers to an arrangement of elements wherein the components so described are configured so as to perform their usual function. Thus, control elements operably linked to a coding sequence are capable of effecting the expression of the coding sequence. The control elements need not be contiguous with the coding sequence, so long as they function to direct the expression thereof. Thus, for example, intervening untranslated yet transcribed sequences can be present between a promoter and the coding sequence and the promoter can still be considered "operably linked" to the coding sequence. Similarly, "control elements compatible with expression in a subject" are those which are capable of effecting the expression of the coding sequence in that subject.

A control element, such as a promoter, "directs the transcription" of a coding sequence in a cell when RNA polymerase will bind the promoter and transcribe the coding sequence into mRNA, which is then translated into the polypeptide encoded by the coding sequence.

A "host cell" is a cell which has been transformed, or is capable of transformation, by an exogenous nucleic acid molecule.

A cell has been "transformed" by exogenous DNA when such exogenous DNA has been introduced inside the cell membrane. Exogenous DNA may or may not be integrated (covalently linked) into chromosomal DNA making up the genome of the cell. In procaryotes and yeasts, for example, the exogenous DNA may be maintained on an episomal element, such as a plasmid. With respect to eucaryotic cells, a stably transformed cell is one in which the exogenous DNA has become integrated into the chromosome so that it is inherited by daughter cells through chromosome replication. This stability is demonstrated by the ability of the eucaryotic cell to establish cell lines or clones comprised of a population of daughter cells containing the exogenous DNA.

As used herein, a "biological sample" refers to a sample of tissue or fluid isolated from a subject, including but not limited to, for example, blood, plasma, serum, fecal matter, urine, bone marrow, bile, spinal fluid, lymph fluid, samples of the skin, external secretions of the skin, respiratory, intestinal, and genitourinary tracts, tears, saliva, milk, blood cells, organs, biopsies and also samples of *in vitro* cell culture constituents including but not limited to conditioned media resulting from the growth of cells and tissues in culture medium, e.g., recombinant cells, and cell components.

As used herein, the terms "label" and "detectable label" refer to a molecule capable of detection, including, but not limited to, radioactive isotopes, fluorescers,

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chemiluminescers, enzymes, enzyme substrates, enzyme cofactors, enzyme inhibitors, chromophores, dyes, metal ions, metal sols, ligands (e.g., biotin or haptens) and the like. The term "fluorescer" refers to a substance or a portion thereof which is capable of exhibiting fluorescence in the detectable range. Particular examples of labels which may be used under the invention include fluorescein, rhodamine, dansyl, umbelliferone, Texas red, luminol, NADPH and α - β -galactosidase.

2. Modes of Carrying Out the Invention

Before describing the present invention in detail, it is to be understood that this invention is not limited to particular formulations or process parameters as such may, of course, vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments of the invention only, and is not intended to be limiting.

Although a number of methods and materials similar or equivalent to those described herein can be used in the practice of the present invention, the preferred materials and methods are described herein.

Central to the present invention is the discovery that the GapC protein is capable of eliciting an immune response in a vertebrate subject. In particular, the genes for the GapC protein in S. dysgalactiae, S. agalactiae, S. uberis, S. parauberis, and S. iniae have been isolated, sequenced and characterized, and the protein sequences for those genes have been deduced therefrom. The complete DNA sequences for those genes and the corresponding amino acid sequences are shown in Figures 1 through 5.

As described in the examples, the full-length *S. dysgalactiae gapC* gene, depicted at nucleotide positions 1-1011, inclusive, of Figures 1A-1B, encodes the full-length *S. dysgalactiae* GapC protein of 336 amino acids, shown as amino acids 1-336, inclusive, of the same figure. *S. dysgalactiae* GapC has a predicted molecular weight of about 36 kDa. (calculated using the Peptide Sort program of the GCG Wisconsin Package, version10, provided by the SeqWeb sequence analysis package, version 1.1 of the Canadian Bioinformatics Resource). Similarly, the *gapC* genes isolated from *S. agalactiae*, *S. uberis*, *S. parauberis* and *S. iniae* are depicted in Figures 2 through 5; each encodes a full-length GapC protein also of 336 amino acids, each also having a predicted molecular weight of about 36 kDa. None of the full-length sequences appear to include a signal peptide or a membrane anchor region.

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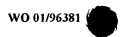


Figures 6 and 7 show an alignment of DNA and amino acid sequences, respectively, showing regions of homology and variability that exist among GapC proteins from various streptococci strains. In particular, several variable regions are located at amino acid positions 62 to 81; 102 to 112; 165 to 172; 248 to 271; and 286 to 305. Such variable regions are typically more amenable to change. Hence, amino acid changes in these regions, such as substitutions, additions and deletions, are likely tolerated.

Figures 8 through 12 show plots of the following for each of the GapC proteins of the present invention: Kyte-Doolittle hydrophathy, averaged over a window of 7; surface probability according to Emini; chain flexibility according to Karplus-Schulz; antigenicity index according to Jameson-Wolf; secondary structure according to Garnier-Osguthorpe-Robson; secondary structure according to Chou-Fasman; and predicted glycosylation sites. Figures 13 through 17 show plots of secondary structure according to Chou-Fasman for each of the GapC proteins of the present invention. One of skill in the art can readily use the information presented in Figures 8 to 17 to determine immunogenic regions in the protein for use in vaccine compositions.

S. dysgalactiae GapC plasmin-binding protein, immunogenic fragments thereof or chimeric proteins including the same, can be provided in subunit vaccine compositions. In addition to use in vaccine compositions, the proteins or antibodies thereto can be used as diagnostic reagents to detect the presence of infection in a vertebrate subject. Similarly, the genes encoding the proteins can be cloned and used to design probes to detect and isolate homologous genes in other bacterial strains. For example, fragments comprising at least about 15-20 nucleotides, more preferably at least about 20-50 nucleotides, and most preferably about 60-100 nucleotides, or any integer between these values, will find use in these embodiments.

The vaccine compositions of the present invention can be used to treat or prevent a wide variety of bacterial infections in vertebrate subjects. For example, vaccine compositions including GapC plasmin-binding proteins from S. dysgalactiae, S. uberis, S. parauberis, S. iniae, and/or group B streptococci (GBS) such as S. agalactiae, can be used to treat streptococcal infections in vertebrate subjects that are caused by these or other species. In particular, S. uberis and S. agalactiae are common bacterial pathogens associated with mastitis in bovine, equine, ovine and goat species. Additionally, group B streptococci, such as S. agalactiae, are known to cause numerous other infections in vertebrates, including septicemia, meningitis, bacteremia, impetigo, arthritis, urinary tract



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infections, abscesses, spontaneous abortion etc. Hence, vaccine compositions containing streptococcal GapC plasmin-binding proteins will find use in treating and/or preventing a wide variety of streptococcal infections.

Similarly, GapC plasmin-binding proteins derived from other bacterial genera such as Staphylococcus, Mycobacterium, Escherichia, Pseudomonas, Nocardia, Pasteurella, Clostridium and Mycoplasma will find use for treating bacterial infections caused by species belonging to those genera. Thus, it is readily apparent that GapC plasmin-binding proteins can be used to treat and /or prevent a wide variety of bacterial infections in numerous species.

The streptococcal GapC plasmin-binding proteins of the present invention can be used in vaccine compositions either alone or in combination with other bacterial, fungal, viral or protozoal antigens. These antigens can be provided separately or even as fusion proteins comprising one or more epitopes of a GapC plasmin-binding protein fused to one or more of these antigens. For example, other immunogenic proteins from S. uberis, such as the CAMP factor, hyaluronic acid capsule, hyaluronidase, R-like protein and plasminogen activator, can be administered with the GapC protein. Additionally, immunogenic proteins from other organisms involved in mastitis, such as from the genera Staphylococcus, Corynebacterium, Pseudomonas, Nocardia, Clostridium, Mycobacterium, Mycoplasma. Pasteurella, Prototheca, other streptococci, coliform bacteria, as well as yeast, can be administered along with the GapC plasmin-binding proteins described herein to provide a broad spectrum of protection. Thus, for example, immunogenic proteins from Staphylococcus aureus, Str. agalactiae, Str. dysgalactiae, Str. zooepidemicus, Corynebacterium pyogenes, Pseudomonas aeruginosa, Nocardia asteroides, Clostridium perfringens, Escherichia coli, Enterobacter aerogenes and Klebsiella spp. can be provided along with the GapC plasmin-binding proteins of the present invention.

Additionally, GapC proteins from different streptococcal species may be used together in the vaccine compositions of the present invention. In this embodiment, the multiple GapC proteins may be provided as fusion proteins or as discrete antigens in the same or different vaccine compositions.

Production of GapC Plasmin-Binding Proteins

The above-described plasmin-binding proteins and active fragments, analogs and chimeric proteins derived from the same, can be produced by variety of methods.

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Specifically, GapC plasmin-binding proteins can be isolated directly from bacteria which express the same. This is generally accomplished by first preparing a crude extract which lacks cellular components and several extraneous proteins. The desired proteins can then be further purified from the cell lysate fraction by, e.g., column chromatography, HPLC, immunoadsorbent techniques or other conventional methods well known in the art.

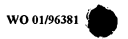
More particularly, techniques for isolating GapC plasmin-binding proteins have been described. For example, the GapC protein of *S. pyogenes* was purified from a crude cell extract by precipitation with ammonium sulfate, followed by two cycles of chromatography through a Mono FPLC column, and single cycles through superose 12 FPLC, and TSK-phenol HPLC columns (Pancholi, V. and Fischetti, VA (1992) *J Exptl. Med* 76:415-426). Another technique involves the use of a NAD⁺-agarose affinity column to purify GapC from lysed protoplasts of *S. pyogenes* strain 64/14 (Winram, SB and Lottenberg, R (1996) *Microbiol.* 142:2311-2320).

Alternatively, the proteins can be recombinantly produced as described herein. As explained above, these recombinant products can take the form of partial protein sequences, full-length sequences, precursor forms that include signal sequences, mature forms without signals, or even fusion proteins (e.g., with an appropriate leader for the recombinant host, or with another subunit antigen sequence for *Streptococcus* or another pathogen).

In one embodiment of the present invention, the GapC proteins are fused to a histidine tag, produced by recombinant means, and are then purified from a cell lysate fraction using affinity chromatography. See Example 1A-E, *infra*.

The GapC plasmin-binding proteins of the present invention can be isolated based on the ability of the protein products to bind plasmin, using plasmin-binding assays as described below. See, e.g., the method described in section F.3. of Example 1, *infra*. Thus, gene libraries can be constructed and the resulting clones used to transform an appropriate host cell. Colonies can be pooled and screened for clones having plasmin-binding activity. Colonies can also be screened using polyclonal serum or monoclonal antibodies to the plasmin-binding protein.

Alternatively, once the amino acid sequences are determined, oligonucleotide probes which contain the codons for a portion of the determined amino acid sequences can be prepared and used to screen genomic or cDNA libraries for genes encoding the subject proteins. The basic strategies for preparing oligonucleotide probes and DNA libraries, as well as their screening by nucleic acid hybridization, are well known to those of ordinary



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skill in the art. See, e.g., DNA Cloning: Vol. 1I, supra; Nucleic Acid Hybridization, supra; Oligonucleotide Synthesis, supra; Sambrook et al., supra. Once a clone from the screened library has been identified by positive hybridization, it can be confirmed by restriction enzyme analysis and DNA sequencing that the particular library insert contains GapC plasmin-binding protein gene or a homolog thereof. The genes can then be further isolated using standard techniques and, if desired, PCR approaches or restriction enzymes employed to delete portions of the full-length sequence.

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Similarly, genes can be isolated directly from bacteria using known techniques, such as phenol extraction and the sequence further manipulated to produce any desired alterations. See, e.g., Sambrook et al., *supra*, for a description of techniques used to obtain and isolate DNA.

Alternatively, DNA sequences encoding the proteins of interest can be prepared synthetically rather than cloned. The DNA sequences can be designed with the appropriate codons for the particular amino acid sequence. In general, one will select preferred codons for the intended host if the sequence will be used for expression. The complete sequence is assembled from overlapping oligonucleotides prepared by standard methods and assembled into a complete coding sequence. See, e.g., Edge (1981) *Nature* 292:756; Nambair et al. (1984) *Science* 223:1299; Jay et al. (1984) *J. Biol. Chem.* 259:6311.

Once coding sequences for the desired proteins have been prepared or isolated, they can be cloned into any suitable vector or replicon. Numerous cloning vectors are known to those of skill in the art, and the selection of an appropriate cloning vector is a matter of choice. Examples of recombinant DNA vectors for cloning and host cells which they can transform include the bacteriophage λ (E. coli), pBR322 (E. coli), pACYC177 (E. coli), pKT230 (gram-negative bacteria), pGV1106 (gram-negative bacteria), pLAFR1 (gram-negative bacteria), pME290 (non-E. coli gram-negative bacteria), pHV14 (E. coli and Bacillus subtilis), pBD9 (Bacillus), pIJ61 (Streptomyces), pUC6 (Streptomyces), YIp5 (Saccharomyces), YCp19 (Saccharomyces) and bovine papilloma virus (mammalian cells). See, Sambrook et al., supra; DNA Cloning, supra; B. Perbal, supra.

The gene can be placed under the control of a promoter, ribosome binding site (for bacterial expression) and, optionally, an operator (collectively referred to herein as "control" elements), so that the DNA sequence encoding the desired protein is transcribed into RNA in the host cell transformed by a vector containing this expression construction. The coding sequence may or may not contain a signal peptide or leader sequence. If a signal sequence

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is included, it can either be the native, homologous sequence, or a heterologous sequence. For example, the signal sequence for *S. dysgalactiae* GapC plasmin-binding protein can be used for secretion thereof, as can a number of other signal sequences, well known in the art. Leader sequences can be removed by the host in post-translational processing. See, e.g., U.S. Patent Nos. 4,431,739; 4,425,437; 4,338,397.

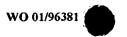
Other regulatory sequences may also be desirable which allow for regulation of expression of the protein sequences relative to the growth of the host cell. Regulatory sequences are known to those of skill in the art, and examples include those which cause the expression of a gene to be turned on or off in response to a chemical or physical stimulus, including the presence of a regulatory compound. Other types of regulatory elements may also be present in the vector, for example, enhancer sequences.

The control sequences and other regulatory sequences may be ligated to the coding sequence prior to insertion into a vector, such as the cloning vectors described above.

Alternatively, the coding sequence can be cloned directly into an expression vector which already contains the control sequences and an appropriate restriction site.

In some cases it may be necessary to modify the coding sequence so that it may be attached to the control sequences with the appropriate orientation; i.e., to maintain the proper reading frame. It may also be desirable to produce mutants or analogs of the GapC plasmin-binding protein. Mutants or analogs may be prepared by the deletion of a portion of the sequence encoding the protein, by insertion of a sequence, and/or by substitution of one or more nucleotides within the sequence. Techniques for modifying nucleotide sequences, such as site-directed mutagenesis, are described in, e.g., Sambrook et al., supra; DNA Cloning, supra; Nucleic Acid Hybridization, supra.

The expression vector is then used to transform an appropriate host cell. A number of mammalian cell lines are known in the art and include immortalized cell lines available from the American Type Culture Collection (ATCC), such as, but not limited to, Chinese hamster ovary (CHO) cells, HeLa cells, baby hamster kidney (BHK) cells, monkey kidney cells (COS), human hepatocellular carcinoma cells (e.g., Hep G2), Madin-Darby bovine kidney ("MDBK") cells, as well as others. Similarly, bacterial hosts such as *E. coli*, *Bacillus subtilis*, and *Streptococcus spp.*, will find use with the present expression constructs. Yeast hosts useful in the present invention include *inter alia*, *Saccharomyces cerevisiae*, *Candida albicans*, *Candida maltosa*, *Hansenula polymorpha*, *Kluyveromyces fragilis*, *Kluyveromyces lactis*, *Pichia guillerimondii*, *Pichia pastoris*, *Schizosaccharomyces*



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pombe and Yarrowia lipolytica. Insect cells for use with baculovirus expression vectors include, inter alia, Aedes aegypti, Autographa californica, Bombyx mori, Drosophila melanogaster, Spodoptera frugiperda, and Trichoplusia ni.

Depending on the expression system and host selected, the proteins of the present invention are produced by culturing host cells transformed by an expression vector described above under conditions whereby the protein of interest is expressed. The protein is then isolated from the host cells and purified. If the expression system secretes the protein into the growth media, the protein can be purified directly from the media. If the protein is not secreted, it is isolated from cell lysates. The selection of the appropriate growth conditions and recovery methods are within the skill of the art.

The proteins of the present invention may also be produced by chemical synthesis such as solid phase peptide synthesis, using known amino acid sequences or amino acid sequences derived from the DNA sequence of the genes of interest. Such methods are known to those skilled in the art. See, e.g., J. M. Stewart and J. D. Young, Solid Phase Peptide Synthesis, 2nd Ed., Pierce Chemical Co., Rockford, IL (1984) and G. Barany and R. B. Merrifield, The Peptides: Analysis, Synthesis, Biology, editors E. Gross and J. Meienhofer, Vol. 2, Academic Press, New York, (1980), pp. 3-254, for solid phase peptide synthesis techniques; and M. Bodansky, Principles of Peptide Synthesis, Springer-Verlag, Berlin (1984) and E. Gross and J. Meienhofer, Eds., The Peptides: Analysis, Synthesis, Biology, supra, Vol. 1, for classical solution synthesis. Chemical synthesis of peptides may be preferable if a small fragment of the antigen in question is capable of raising an immunological response in the subject of interest.

The GapC plasmin-binding proteins of the present invention, or their fragments, can be used to produce antibodies, both polyclonal and monoclonal. If polyclonal antibodies are desired, a selected mammal, (e.g., mouse, rabbit, goat, horse, etc.) is immunized with an antigen of the present invention, or its fragment, or a mutated antigen. Serum from the immunized animal is collected and treated according to known procedures. See, e.g., Jurgens et al. (1985) *J. Chrom.* 348:363-370. If serum containing polyclonal antibodies is used, the polyclonal antibodies can be purified by immunoaffinity chromatography, using known procedures.

Monoclonal antibodies to the GapC plasmin-binding proteins and to the fragments thereof, can also be readily produced by one skilled in the art. The general methodology for making monoclonal antibodies by using hybridoma technology is well known. Immortal

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antibody-producing cell lines can be created by cell fusion, and also by other techniques such as direct transformation of B lymphocytes with oncogenic DNA, or transfection with Epstein-Barr virus. See, e.g., M. Schreier et al., *Hybridoma Techniques* (1980); Hammerling *et al.*, *Monoclonal Antibodies and T-cell Hybridomas* (1981); Kennett et al., *Monoclonal Antibodies* (1980); see also U.S. Patent Nos. 4,341,761; 4,399,121; 4,427,783; 4,444,887; 4,452,570; 4,466,917; 4,472,500, 4,491,632; and 4,493,890. Panels of monoclonal antibodies produced against the GapC plasmin-binding proteins, or fragments thereof, can be screened for various properties; i.e., for isotype, epitope, affinity, etc. Monoclonal antibodies are useful in purification, using immunoaffinity techniques, of the individual antigens which they are directed against. Both polyclonal and monoclonal antibodies can also be used for passive immunization or can be combined with subunit vaccine preparations to enhance the immune response. Polyclonal and monoclonal antibodies are also useful for diagnostic purposes.

15 Vaccine Formulations and Administration

The GapC plasmin-binding proteins of the present invention can be formulated into vaccine compositions, either alone or in combination with other antigens, for use in immunizing subjects as described below. Methods of preparing such formulations are described in, e.g., Remington's Pharmaceutical Sciences, Mack Publishing Company, Easton, Pennsylvania, 18 Edition, 1990. Typically, the vaccines of the present invention are prepared as injectables, either as liquid solutions or suspensions. Solid forms suitable for solution in or suspension in liquid vehicles prior to injection may also be prepared. The preparation may also be emulsified or the active ingredient encapsulated in liposome vehicles. The active immunogenic ingredient is generally mixed with a compatible pharmaceutical vehicle, such as, for example, water, saline, dextrose, glycerol, ethanol, or the like, and combinations thereof. In addition, if desired, the vehicle may contain minor amounts of auxiliary substances such as wetting or emulsifying agents and pH buffering agents.

Adjuvants which enhance the effectiveness of the vaccine may also be added to the formulation. Adjuvants may include for example, muramyl dipeptides, avridine, aluminum hydroxide, dimethyldioctadecyl ammonium bromide (DDA), oils, oil-in-water emulsions, saponins, cytokines, and other substances known in the art.

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The GapC plasmin-binding protein may be linked to a carrier in order to increase the immunogenicity thereof. Suitable carriers include large, slowly metabolized macromolecules such as proteins, including serum albumins, keyhole limpet hemocyanin, immunoglobulin molecules, thyroglobulin, ovalbumin, and other proteins well known to those skilled in the art; polysaccharides, such as sepharose, agarose, cellulose, cellulose beads and the like; polymeric amino acids such as polyglutamic acid, polylysine, and the like; amino acid copolymers; and inactive virus particles.

The GapC plasmin-binding proteins may be used in their native form or their functional group content may be modified by, for example, succinylation of lysine residues or reaction with Cys-thiolactone. A sulfhydryl group may also be incorporated into the carrier (or antigen) by, for example, reaction of amino functions with 2-iminothiolane or the N-hydroxysuccinimide ester of 3-(4-dithiopyridyl propionate. Suitable carriers may also be modified to incorporate spacer arms (such as hexamethylene diamine or other bifunctional molecules of similar size) for attachment of peptides.

Other suitable carriers for the GapC plasmin-binding proteins of the present invention include VP6 polypeptides of rotaviruses, or functional fragments thereof, as disclosed in U.S. Patent No. 5,071,651. Also useful is a fusion product of a viral protein and the subject immunogens made by methods disclosed in U.S. Patent No. 4,722,840. Still other suitable carriers include cells, such as lymphocytes, since presentation in this form mimics the natural mode of presentation in the subject, which gives rise to the immunized state. Alternatively, the proteins of the present invention may be coupled to erythrocytes, preferably the subject's own erythrocytes. Methods of coupling peptides to proteins or cells are known to those of skill in the art.

Furthermore, the GapC plasmin-binding proteins (or complexes thereof) may be formulated into vaccine compositions in either neutral or salt forms. Pharmaceutically acceptable salts include the acid addition salts (formed with the free amino groups of the active polypeptides) and which are formed with inorganic acids such as, for example, hydrochloric or phosphoric acids, or such organic acids as acetic, oxalic, tartaric, mandelic, and the like. Salts formed from free carboxyl groups may also be derived from inorganic bases such as, for example, sodium, potassium, ammonium, calcium, or ferric hydroxides, and such organic bases as isopropylamine, trimethylamine, 2-ethylamino ethanol, histidine, procaine, and the like.

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Vaccine formulations will contain a "therapeutically effective amount" of the active ingredient, that is, an amount capable of eliciting an immune response in a subject to which the composition is administered. In the treatment and prevention of mastitis, for example, a "therapeutically effective amount" would preferably be an amount that enhances resistance of the mammary gland to new infection and/or reduces the clinical severity of the disease. Such protection will be demonstrated by either a reduction or lack of symptoms normally displayed by an infected host, a quicker recovery time and/or a lowered somatic cell count in milk from the infected quarter. For example, the ability of the composition to retain or bring the somatic cell count (SCC) in milk below about 500,000 cells per ml, the threshold value set by the International Dairy Federation, above which, animals are considered to have clinical mastitis, will be indicative of a therapeutic effect.

The exact amount is readily determined by one skilled in the art using standard tests. The GapC plasmin-binding protein concentration will typically range from about 1% to about 95% (w/w) of the composition, or even higher or lower if appropriate. With the present vaccine formulations, 5 to 500 µg of active ingredient per ml of injected solution should be adequate to raise an immunological response when a dose of 1 to 3 ml per animal is administered.

To immunize a subject, the vaccine is generally administered parenterally, usually by intramuscular injection. Other modes of administration, however, such as subcutaneous, intraperitoneal and intravenous injection, are also acceptable. The quantity to be administered depends on the animal to be treated, the capacity of the animal's immune system to synthesize antibodies, and the degree of protection desired. Effective dosages can be readily established by one of ordinary skill in the art through routine trials establishing dose response curves. The subject is immunized by administration of the vaccine in at least one dose, and preferably two doses. Moreover, the animal may be administered as many doses as is required to maintain a state of immunity to infection.

Additional vaccine formulations which are suitable for other modes of administration include suppositories and, in some cases, aerosol, intranasal, oral formulations, and sustained release formulations. For suppositories, the vehicle composition will include traditional binders and carriers, such as, polyalkaline glycols, or triglycerides. Such suppositories may be formed from mixtures containing the active ingredient in the range of about 0.5% to about 10% (w/w), preferably about 1% to about 2%. Oral vehicles include such normally employed excipients as, for example,

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pharmaceutical grades of mannitol, lactose, starch, magnesium, stearate, sodium saccharin cellulose, magnesium carbonate, and the like. These oral vaccine compositions may be taken in the form of solutions, suspensions, tablets, pills, capsules, sustained release formulations, or powders, and contain from about 10% to about 95% of the active ingredient, preferably about 25% to about 70%.

Intranasal formulations will usually include vehicles that neither cause irritation to the nasal mucosa nor significantly disturb ciliary function. Diluents such as water, aqueous saline or other known substances can be employed with the subject invention. The nasal formulations may also contain preservatives such as, but not limited to, chlorobutanol and benzalkonium chloride. A surfactant may be present to enhance absorption of the subject proteins by the nasal mucosa.

Controlled or sustained release formulations are made by incorporating the protein into carriers or vehicles such as liposomes, nonresorbable impermeable polymers such as ethylenevinyl acetate copolymers and Hytrel® copolymers, swellable polymers such as hydrogels, or resorbable polymers such as collagen and certain polyacids or polyesters such as those used to make resorbable sutures. The GapC plasmin-binding proteins can also be delivered using implanted mini-pumps, well known in the art.

The GapC plasmin-binding proteins of the instant invention can also be administered via a carrier virus which expresses the same. Carrier viruses which will find use with the instant invention include but are not limited to the vaccinia and other pox viruses, adenovirus, and herpes virus. By way of example, vaccinia virus recombinants expressing the novel proteins can be constructed as follows. The DNA encoding the particular protein is first inserted into an appropriate vector so that it is adjacent to a vaccinia promoter and flanking vaccinia DNA sequences, such as the sequence encoding thymidine kinase (TK). This vector is then used to transfect cells which are simultaneously infected with vaccinia. Homologous recombination serves to insert the vaccinia promoter plus the gene encoding the instant protein into the viral genome. The resulting TK recombinant can be selected by culturing the cells in the presence of 5-bromodeoxyuridine and picking viral plaques resistant thereto.

An alternative route of administration involves gene therapy or nucleic acid immunization. Thus, nucleotide sequences (and accompanying regulatory elements) encoding the subject GapC plasmin-binding proteins can be administered directly to a subject for *in vivo* translation thereof. Alternatively, gene transfer can be accomplished by

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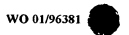


transfecting the subject's cells or tissues ex vivo and reintroducing the transformed material into the host. DNA can be directly introduced into the host organism, i.e., by injection (see International Publication No. WO/90/11092; and Wolff et al. (1990) Science 247:1465-1468). Liposome-mediated gene transfer can also be accomplished using known methods. Sce, e.g., Hazinski et al. (1991) Am. J. Respir. Cell Mol. Biol. 4:206-209; Brigham et al. (1989) Am. J. Med. Sci. 298:278-281; Canonico et al. (1991) Clin. Res. 39:219A; and Nabel et al. (1990) Science 249:1285-1288. Targeting agents, such as antibodies directed against surface antigens expressed on specific cell types, can be covalently conjugated to the liposomal surface so that the nucleic acid can be delivered to specific tissues and cells susceptible to infection.

Diagnostic Assays

As explained above, the GapC plasmin-binding proteins of the present invention may also be used as diagnostics to detect the presence of reactive antibodies of streptococcus, for example *S. dysgalactiae*, in a biological sample in order to determine the presence of streptococcus infection. For example, the presence of antibodies reactive with GapC plasmin-binding proteins can be detected using standard electrophoretic and immunodiagnostic techniques, including immunoassays such as competition, direct reaction, or sandwich type assays. Such assays include, but are not limited to, Western blots; agglutination tests; enzyme-labeled and mediated immunoassays, such as ELISAs; biotin/avidin type assays; radioimmunoassays; immunoelectrophoresis; immunoprecipitation, etc. The reactions generally include revealing labels such as fluorescent, chemiluminescent, radioactive, enzymatic labels or dye molecules, or other methods for detecting the formation of a complex between the antigen and the antibody or antibodies reacted therewith.

The aforementioned assays generally involve separation of unbound antibody in a liquid phase from a solid phase support to which antigen-antibody complexes are bound. Solid supports which can be used in the practice of the invention include substrates such as nitrocellulose (e.g., in membrane or microtiter well form); polyvinylchloride (e.g., sheets or microtiter wells); polystyrene latex (e.g., beads or microtiter plates); polyvinylidine fluoride; diazotized paper; nylon membranes; activated beads, magnetically responsive beads, and the like.



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Typically, a solid support is first reacted with a solid phase component (e.g., one or more GapC plasmin-binding proteins) under suitable binding conditions such that the component is sufficiently immobilized to the support. Sometimes, immobilization of the antigen to the support can be enhanced by first coupling the antigen to a protein with better binding properties. Suitable coupling proteins include, but are not limited to, macromolecules such as serum albumins including bovine serum albumin (BSA), keyhole limpet hemocyanin, immunoglobulin molecules, thyroglobulin, ovalbumin, and other proteins well known to those skilled in the art. Other molecules that can be used to bind the antigens to the support include polysaccharides, polylactic acids, polyglycolic acids, polymeric amino acids, amino acid copolymers, and the like. Such molecules and methods of coupling these molecules to the antigens, are well known to those of ordinary skill in the art. See, e.g., Brinkley, M.A. Bioconjugate Chem. (1992) 3:2-13; Hashida et al., J. Appl. Biochem. (1984) 6:56-63; and Anjaneyulu and Staros, International J. of Peptide and Protein Res. (1987) 30:117-124.

After reacting the solid support with the solid phase component, any non-immobilized solid-phase components are removed from the support by washing, and the support-bound component is then contacted with a biological sample suspected of containing ligand moieties (e.g., antibodies toward the immobilized antigens) under suitable binding conditions. After washing to remove any non-bound ligand, a secondary binder moiety is added under suitable binding conditions, wherein the secondary binder is capable of associating selectively with the bound ligand. The presence of the secondary binder can then be detected using techniques well known in the art.

More particularly, an ELISA method can be used, wherein the wells of a microtiter plate are coated with a GapC plasmin-binding protein. A biological sample containing or suspected of containing anti-GapC plasmin-binding protein immunoglobulin molecules is then added to the coated wells. After a period of incubation sufficient to allow antibody binding to the immobilized antigen, the plate(s) can be washed to remove unbound moieties and a detectably labeled secondary binding molecule added. The secondary binding molecule is allowed to react with any captured sample antibodies, the plate washed and the presence of the secondary binding molecule detected using methods well known in the art.

Thus, in one particular embodiment, the presence of bound anti-GapC plasminbinding antigen ligands from a biological sample can be readily detected using a secondary binder comprising an antibody directed against the antibody ligands. A number of anti-

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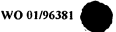


bovine immunoglobulin (Ig) molecules are known in the art which can be readily conjugated to a detectable enzyme label, such as horseradish peroxidase, alkaline phosphatase or urease, using methods known to those of skill in the art. An appropriate enzyme substrate is then used to generate a detectable signal. In other related embodiments, competitive-type ELISA techniques can be practiced using methods known to those skilled in the art.

Assays can also be conducted in solution, such that the GapC plasmin-binding proteins and antibodies specific for those proteins form complexes under precipitating conditions. In one particular embodiment, GapC plasmin-binding proteins can be attached to a solid phase particle (e.g., an agarose bead or the like) using coupling techniques known in the art, such as by direct chemical or indirect coupling. The antigen-coated particle is then contacted under suitable binding conditions with a biological sample suspected of containing antibodies for the GapC plasmin-binding proteins. Cross-linking between bound antibodies causes the formation of particle-antigen-antibody complex aggregates which can be precipitated and separated from the sample using washing and/or centrifugation. The reaction mixture can be analyzed to determine the presence or absence of antibody-antigen complexes using any of a number of standard methods, such as those immunodiagnostic methods described above.

In yet a further embodiment, an immunoaffinity matrix can be provided, wherein a polyclonal population of antibodies from a biological sample suspected of containing anti-GapC plasmin-binding molecules is immobilized to a substrate. In this regard, an initial affinity purification of the sample can be carried out using immobilized antigens. The resultant sample preparation will thus only contain anti-streptococcus moieties, avoiding potential nonspecific binding properties in the affinity support. A number of methods of immobilizing immunoglobulins (either intact or in specific fragments) at high yield and good retention of antigen binding activity are known in the art. Not being limited by any particular method, immobilized protein A or protein G can be used to immobilize immunoglobulins.

Accordingly, once the immunoglobulin molecules have been immobilized to provide an immunoaffinity matrix, labeled GapC plasmin-binding proteins are contacted with the bound antibodies under suitable binding conditions. After any non-specifically bound antigen has been washed from the immunoaffinity support, the presence of bound antigen can be determined by assaying for label using methods known in the art.



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Additionally, antibodies raised to the GapC plasmin-binding proteins, rather than the GapC plasmin-binding proteins themselves, can be used in the above-described assays in order to detect the presence of antibodies to the proteins in a given sample. These assays are performed essentially as described above and are well known to those of skill in the art.

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The above-described assay reagents, including the GapC plasmin-binding proteins, or antibodies thereto, can be provided in kits, with suitable instructions and other necessary reagents, in order to conduct immunoassays as described above. The kit can also contain, depending on the particular immunoassay used, suitable labels and other packaged reagents and materials (i.e. wash buffers and the like). Standard immunoassays, such as those described above, can be conducted using these kits.

Deposits of Strains Useful in Practicing the Invention

A deposit of biologically pure cultures of the following strains was made with the American Type Culture Collection, 10801 University Boulevard, Manassas, Virginia, under the provisions of the Budapest Treaty. The accession number indicated was assigned after successful viability testing, and the requisite fees were paid. The designated deposits will be maintained for a period of thirty (30) years from the date of deposit, or for five (5) years after the last request for the deposit, whichever is longer. Should a culture become nonviable or be inadvertently destroyed, or, in the case of plasmid-containing strains, lose its plasmid, it will be replaced with a viable culture(s) of the same taxonomic description.

Should there be a discrepancy between the sequence presented in the present application and the sequence of the gene of interest in the deposited plasmid due to routine sequencing errors, the sequence in the deposited plasmid controls.

25	Bacterial Strain	Plasmid	Gene	Deposit Date	ATCC No.
	E. coli BL21 DE3	pET15bgapC	gapC (S. dysgalactiae)	May 31, 2000	PTA-1976
30	E. coli BL21 DE3	pMF521c	gapC (S. agalactiae)	May 31, 2000	PTA-1975
	E. coli BL21 DE3	pMF521a	gapC (S. uberis)	May 31, 2000	PTA-1973

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E. coli pMF521e BL21 DE3

gapC (S. iniae) May 31, 2000

PTA-1972

3. Experimental ...

Below are examples of specific embodiments for carrying out the present invention. The examples are offered for illustrative purposes only, and are not intended to limit the scope of the present invention in any way.

Efforts have been made to ensure accuracy with respect to numbers used (e.g., amounts, temperatures, etc.), but some experimental error and deviation should, of course, be allowed for.

Materials and Methods

Enzymes were purchased from commercial sources, and used according to the manufacturers' directions.

In the isolation of DNA fragments, except where noted, all DNA manipulations were done according to standard procedures. See, Sambrook et al., *supra*. Restriction enzymes, T₄ DNA ligase, *E. coli*, DNA polymerase 1I, Klenow fragment, and other biological reagents can be purchased from commercial suppliers and used according to the manufacturers' directions. Double stranded DNA fragments were separated on agarose gels.

Sources for chemical reagents generally include Sigma Chemical Company, St. Louis, MO; Alrich, Milwaukee, WI; Roche Molecular Biochemicals, Indianapolis, IN.

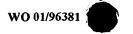
EXAMPLE 1

Preparation, Amplification, Sequencing, Expression, Purification and Characterization of

the S. dysgalactiae GapC Plasmin Binding Protein

A. Preparation of S. dysgalactiae Chromosomal DNA

A clinical *S. dysgalactiae* isolate from a case of bovine mastitis (ATCC Accession No. ATCC43078) was obtained from the American Type Culture Collection (10801 University Boulevard, Manassas, VA 20110-2209), and was used as a source of DNA. The organism was routinely grown on TSA sheep blood agar plates (PML Microbiologicals, Mississauga, Ontario) at 37° C for 18 hours, or in Todd-Hewitt broth (Oxoid Ltd., Hampshire, England) supplemented with 0.3% yeast extract (THB-YE) at 37° C, 5% CO₂.



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Chromosomal DNA was prepared from *S. dysgalactiae* grown in 100 ml of THB-YE supplemented with 20 mM glycine for approximately 6 hours, until an A₆₀₀ of 0.8 to 1.0 was reached. Cells were harvested and re-suspended in 50 mM EDTA, 50 mM Tris-HCl, 0.5% Tween-20® (Sigma, St. Louis, MO) and supplemented with RNase A (200 mg/ml), proteinase K (20 mg/ml), lysozyme (100 mg/ml) and mutanolysin (100 mg/ml). (SIGMA, St. Louis, MO). Following bacterial lysis for 30 minutes at 37° C with vigorous shaking, guanidine hydrochloride and Tween-2®, pH 5.5, were mixed with the lysate to give a final concentration of 0.8 M and 5%, respectively. This mixture was incubated at 50° C for 30 minutes. The chromosomal DNA was then purified using a Qiagen genomic-tip 100g (Qiagen, Santa Clarita, CA) and precipitated using 0.7 volumes of isopropanol. The resulting pellet was washed in 70% ethanol and re-suspended in 0.5 ml 10 mM Tris-HCl, pH 8.8.

B. Amplification and Cloning of the S. dysgalactiae gapC Gene

The gapC gene was amplified by PCR (See Mullis et al., U.S. Patent No. 4,683,195; Mullis, U.S. Patent No. 4,683,202;). The forward primer, gapC1, contained an Nde1 restriction (SEQ ID NO:1, shown in Table 1) and the reverse primer, gapC1r, contained a BamHI site (SEQ ID NO:2, shown in Table 1). In the preceding primer sequences, depicted in Table 1, underlining denotes nucleotides added to the original sequence, and bolding indicates the location of restriction endonuclease recognition sites.

PCR was carried out using Vent DNA polymerase (New England Biolabs, Mississauga, ON, Canada). 0.7 μg of S. dysgalactiae chromosomal DNA was incubated in a reaction mixture containing 1μM of each of the preceding primers, 200 μM each of dATP, dTTP, dCTP and dGTP, 3mM MgSO₄, 1x concentration of Thermopol buffer (New England Biolabs, Mississauga, ON, Canada) and 2 units Vent DNA polymerase. This mixture was incubated for 3 amplification cycles of 1 minute at 94°C, 3 minutes at 50°C and 1 minute, 10 seconds at 72°C, then for 27 amplification cycles at 15 seconds at 95°C, 30 seconds at 55°C, and 1 minute at 72°C, and finally for 1 cycle of 5 minutes at 72°C.



TABLE 1

Sequence Identification Numbers and

Corresponding Nucleotide and Amino Acid Sequences

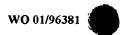
SEQ ID NO.	Name	Sequence		
1	Primer gapC1	5'- <u>GG CGG CGG CAT</u> ATG GTA GTT AAA GTT GGT ATT AAC GG -3'		
2	Primer gapC1r	5'- <u>GC GGA TCC</u> TTA TTT AGC GAT TTT TGC AAA GTA CTC -3'		
3	Streptococcus dysgalactiae gapC gene	(see Figure 1)		
4	Streptococcus dysgalactiae GapC protein			
5	Streptococcus agalactiae gapC gene	(see Figure 2)		
6	Streptococcus agalactiae GapC protein			
7	Streptococcus uberis gapC gene	(see Figure 3)		
8	Streptococcus uberis GapC protein			
9	Streptococcus parauberis gapC gene			
10	Streptococcus parauberis GapC protein	(see Figure 4)		
11	Streptococcus iniae gapC gene	(see Figure 5)		
12	Streptococcus iniae GapC protein			

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The gapC PCR product was cloned into the expression vector pET15B (Novagen, Madison, WI) which had been digested with BamHI and NdeI. Cloning of the PCR product into this site results in the addition of an in-frame coding sequence for a hexahistidyl tag to the gapC coding sequence. Subsequent expression yields a full-length protein with an attached histidine tag, which permits purification of the protein under non-denaturing conditions using metal chelate chromatography.

This construct was used to transform *E. coli* BL21 DE3 (Life Technologies, Gaithersburg, MD). This transformed strain was designated BL21 DE3 (pET15bgapC) (ATCC No:).

C. <u>Isolation of Chromosomal DNA and Amplification and Cloning of the gapC Gene</u> from S. agalactiae, S. uberis, S. parauberis and S. iniae

The gapC gene were prepared from other isolates essentially as described above. Chromosomal DNA from S. agalactiae, S. uberis, and S. parauberis was isolated from strains obtained from the American Type Culture Collection (10801 University Boulevard, Manassas, VA 20110-2209; strains designated ATCC 27541, 9927, and 13386, respectively). Chromosomal DNA from S. iniae was isolated from a strain designated 9117 obtained from Mount Sinai Hospital, University of Toronto.

The primers used to amplify the gapC genes from the *Streptococcus* strains listed above were the same as those used in the case of *S. dysgalactiae*, i.e., primer gapC1 (SEQ ID NO:1) and primer gapC1r (SEQ ID NO:2).

After amplification, the PCR product in each case was cloned into pPCR-Script,

using the cloning protocol described in the PCR-Script Amp cloning Kit (Stratagene, La

Jolla, CA). The PCR product insert was then excised using NdeI and BamHI and re-cloned
into those sites in pE15b using conventional cloning protocols (See e.g., Sambrook et al.,

supra.). The plasmids containing the S. agalactiae, S. uberis, S. parauberis and S. iniae
were designated pMF521c (ATCC No:), pMF521a (ATCC No:), pMF521d (ATCC No:),

and pMF521e (ATCC No:), respectively.

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<u>Nucleotide Sequence of the gapC Gene and Deduced Amino Acid Sequences</u> Sequences homologous to gapC of S. equisimilis homolog (Gase, et al. (1996) European J. of Biochem. 239:42-51) were originally identified while sequencing a linked but unrelated gene of S. dysgalactiae. To obtain the complete sequence of the S. dysgalactiae gapC gene, PCR was employed using the primers described above, i.e., primer gapC1 and primer gapC1r.

The sequence was determined using fluorescence tag terminators on an ABI 373 DNA automatic sequencer (Applied Biosystems, Emeryville, CA) at the Plant Biotechnology Institute (PBI, Saskatoon, Canada).

both 1 depicts the coding sequence of the gapC gene from S. dysgalactiae (DysGapC) (SEQ ID NO:3) and the deduced amino acid sequence (SEQ ID NO:4).

The sequences of the GapC proteins isolated form S. agalactiae, S. uberis, S. parauberis, and S. iniae were determined by the same method.

Figures 2 through 5 depict both the nucleotide sequences and the predicted amino acid sequences for the *S. dysgalactiae* GapC protein (DysgalGapC) (SEQ ID NO:3 and SEQ ID NO:4), as well as for the GapC proteins of S. *agalactiae* (AgalGapC) (SEQ ID NO:5 and SEQ ID NO:6), *S. uberis* (UberGapC) (SEQ ID NO:7 and SEQ ID NO:8), *S. parauberis* (PUberGapC) (SEQ ID NO:9 and SEQ ID NO:10), and *S. iniae* (IniaeGapC) (SEQ ID NO:11 and SEQ ID NO:12), respectively.

The S. dysgalactiae GapC protein gene depicted in Figures 1A-1B codes for a 336 amino acid protein which does not appear to contain either a signal sequence or membrane anchor domain. A search of the GenBank database using the BLASTX program revealed that the open reading frame was 95.5% homologous to GapC of S. equisimilis (GenBank Accession No. X97788) and 99.4% homologous to GapC of S. pyogenes (GenBank Accession No. M95569). The predicted amino acid sequence of the GapC protein also exhibited 43% amino acid identity to bovine glyceraldehyde-3-phosphate dehydrogenase (GenBank Accession No. U85042).

Similarly, for the S. agalactiae, S. uberis, S. parauberis and S. iniae GapC protein sequences, neither signal sequences nor membrane anchor domains appear to be present.

Sequence homologies are tabulated in Table 2.

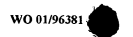


TABLE 2 Sequence Homologies Between Various GapC Protein Sequences

- 	S. equisimilis	S. pyogenes	Bovine GAPDH
S. dysgalactiae	95.5%	94.4%	43%
S. agalactiae	87.02%	91.07%	
S. parauberis	86.31%	90.77%	(not determined)
S. uberis	88.39%	92.26%	
S. iniae	86.31%	89.88%	

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E. Expression and Purification of the Recombinant S. dysgalactiae GapC Plasmin Binding Protein

The Hexahistidyl-tagged GapC protein was expressed and purified under nondenaturing conditions using metal chelate (Ni-NTA) affinity chromatography.

 $E.\ coli$ BL21 DE3 containing the recombinant plasmid was grown in Luria Broth, containing 100 μg/ml ampicillin to an A_{600} of approximately 0.5. Expression of the GapC protein was then induced by the addition of 1 mM isopropyl- β ,D-thiogalactoside (IPTG) (Sigma, St. Louis, MO]. Following three hours incubation at 37°C, cells were harvested, washed in column buffer (50 mM sodium phosphate buffer, pH8.0, 0.3 M NaCl, 10 mM imidazole) and lysed by sonication.

Approximately 40% of the recombinant protein was in the soluble fraction of the cell sonicate with a yield of approximately 50 mg of the recombinant protein per litre of culture volume, determined with a *DC* Protein Assay Kit (Bio-Rad Laboratories, Mississauga, ON, Canada) using bovine serum albumin (Pierce, Rockford, IL) as a standard.

The lysate was cleared by centrifugation and the soluble fraction was applied to a Ni-NTA column (Qiagen), which was subsequently washed with 10 column volumes of column buffer (as above, except containing 20 mM imidazole). The Hexahistidyl-tagged GapC was eluted using column buffer (as above, except containing 250 mM imidazole), yielding a homogenous protein fraction having a GapC concentration of 10-15 mg/ml. That fraction was dialyzed against 2000 volumes of PBSA (136mM sodium chloride, 2.6mM

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potassium chloride, 8.1mM sodium phosphate dibasic, 1.46 mM potassium phosphate monobasic).

F. Expression and Purification of Recombinant GapC Protein from S. agalactiae, S. uberis, S. parauberis, and S. iniae

Expression and purification of the recombinant proteins from these *streptococcus* species is accomplished by the same methods described in Example 1E, above. The transformed bacterial strains used to express the *S. agalactiae*, *S. uberis*, *S. parauberis* and *S. iniae* recombinant GapC proteins were designated BL21 DE3 (pMF521c) (ATCC No:), BL21 DE3 (pMF521a) (ATCC No:), BL21 DE3 (pMF521b) (ATCC No:), and BL21 DE3 (pMF521e) (ATCC No:), respectively.

G. Characterization of the Recombinant S. dysgalactiae GapC Protein

1. <u>SDS-Page Analysis</u>

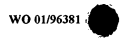
SDS-polyacrylamide gel electrophoresis was performed on a sample of the eluted protein using the method described by Laemli (Laemli, U.K. (1970) *Nature* 227:680-685). The results are presented in Figure 18. In the figure: lane 1, molecular weight markers (20.5 to 103 kDa range; BioRad Laboratories, Emeryville, CA); lane 2, soluble recombinant *S. dysgalactiae* GapC protein purified by Ni-NTA affinity chromatography.

These results demonstrate that purification by affinity chromatography on a Ni-NTA column yielded a homogenous protein fraction.

2. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) activity of recombinant GapC and S. dysgalactiae whole cells

GAPDH catalyzes the oxidative phosphorylation of D-glyceraldehyde-3-phosphate to 1, 3-diphosphoglycerate in the presence of NAD⁺ and inorganic phosphate. The high degree of homology of GapC to streptococcal glyceraldehyde-3-phosphate dehydrogenase suggested that GapC may exhibit this enzyme activity.

The GAPDH activity of S. dysgalactiae whole cells (10¹⁰ CFU) and the recombinant GapC protein was determined by measuring the reduction of NAD⁺ to NADH. The assay buffer was composed of 40 mM triethanolamine, 50 mM Na₂ HPO₄ and 5 mM EDTA, pH 6.8. S. dysgalactiae cells or 5 mg of purified recombinant protein were incubated in assay buffer containing 7 ml glyceraldehyde-3-phosphate (49 mg/ml; Sigma Chemical Company),



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75 microliters NAD⁺ (15 mM; Sigma Chemical Company) in a final volume of 1 ml. Negative controls consisted of samples which did not contain glyeraldehyde-3-phosphate or the recombinant GapC molecule/S. dysgalactiae cells. The reduction of NAD⁺ to NADH was monitored spectrophotometrically at an Absorbance of 340 nanometers.

The results indicated that both the recombinant protein as well as intact wild-type S. dysgalactiea cells had enzymatic activity (not shown). Furthermore, when S. dysgalactiae cells were treated with Trypsin to digest surface proteins, GAPDH activity disappeared. Thus, the enzymatic activity observed for the intact wild type cells was not due to intracellular GAPDH.

This data suggests that the GapC protein is localized on the cell surface despite the apparent lack of either a signal sequence or a membrane anchoring region in either the nucleotide or amino acid.

3. <u>Plasmin-Binding Activity of Recombinant S. dysgalactiae GapC Plasmin Binding Protein and S. dysgalactiae whole cells.</u>

A microplate assay was used to determine if the recombinant GapC protein was capable of binding bovine plasmin, and if so whether the bound plasmin was in an enzymatically active form.

Ninety-six-well microtiter plates were coated with 5 mg of purified recombinant GapC protein, washed 3 times with 0.1% gelatin-PBSA with 0.05% TWEEN-20 (PBSGT). The wells were blocked for one hour at 37° C in the same buffer, washed, and incubated with 200 ml of bovine plasmin (0.25mg/ml; Boehringer Mannheim, Indianapolis, IN) for 1 hr at 37° C. The wells were then washed 8 times with PBSGT. 200 ml of the synthetic substrate chromazine-PL (Tos-Gly-Pro-Lys-4-NA, 0.3 mg/ml) were added to the wells and incubated at 37° C for one hour. The presence of associated plasmin activity was determined by measuring the level of paranitroanalide (4-nitraniline) released into the supernatant and detected based on an Absorbance of 405 nanometers. A similar procedure was used to measure plasmin-binding activity of *S. dysgalactiae* whole cells, with the exception that 10¹⁰ cells were washed with PBSGT and re-suspended in 400 µl chromazine-PL (0.3mg/ml) and incubated for 1 hour at 37° C.

The results, shown in Figure 19, demonstrate that the purified recombinant protein was capable of binding enymatically active bovine plasmin. Likewise, when S.



dysgalactiae whole cells were utilized, similar results were obtained. In the figure, the data represents the mean of three individual assays.

Thus, the plasmin-receptor is located on the surface of *S. dysgalactiae* and the purified protein retains biological activity.

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EXAMPLE 2

Immunization with S. dysgalactiae GapC and experimental infection of cattle

Vaccines were formulated in such a fashion that they contained 50 mg/ml of affinity purified recombinant GapC in the oil-based adjuvant VSA3 (VIDO, Saskatoon, Saskatchewan, Canada). VSA3 is a combination of Emulsigen PlusTM (MVP Laboratories, Ralston, Nebraska) and Dimethyldioctadecyl ammonium bromide (Kodak, Rochester, New York). The affinity-purified recombinant GapC protein used for the vaccine preparation is shown in Figure 18.

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Twenty-four non-lactating Holsteins with no history of *S. dysgalactiae* infection were obtained from various farms in Saskatchewan, Canada. One week prior to vaccination, all animals were treated with Cepha-dryTM (Ayerst Laboratories, Montreal, Canada) (300 mg per quarter), in order to clear any infection of the udders prior to the vaccination step.

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Groups of 8 animals were immunized subcutaneously with two doses of vaccines containing S. dysgalactiae GapC, Mig (an Fc receptor protein isolated from S. dysgalactiae which was evaluated simultaneously), or a placebo with a three-week interval between immunizations. Two weeks following the second immunization, animals were exposed to 650 colony forming units of S. dysgalactiae delivered into three quarters with an udder infusion cannula. The fourth quarter on each animal served as an un-infective control.

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All animals were examined daily for clinical signs of disease and samples from all udder quarters were collected on each day. Samples were observed for consistency and somatic cell counts as well as bacterial numbers were determined.

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EXAMPLE 3

Determination of GapC-specific Antibodies

GapC-specific antibodies in bovine serum were measured using an enzyme-linked immunosorbent assay (ELISA). Briefly, microtiter plates (NUNC, Naperville, Illinois)

were coated by adding 1 microgram per well purified recombinant antigen in 50mM sodium carbonate buffer, pH 9.6, and incubated overnight at 4° C. The liquid was removed and the wells were blocked with 3% bovine serum albumin for 1 hr at 37° C. Serial dilutions of bovine serum (from 1 in 4 to 1 in 6,400) were then added to the wells and incubated for 2 hours at room temperature. The wells were aspirated, washed and incubated with 100 ml of alkaline phosphatase-conjugated goat anti-bovine IgG (Kirkgaard & Perry Laboratories Inc.,Gaithersburg, MD) for 1 hr at room temperature. The wells were washed again, and $100 \mu l$ of p-nitrophenol phosphate (SIGMA, St. Louis, MO) was added as a substrate to detect alkaline phosphatase activity. The absorbance at 405 nanometers was recorded following 1 hr incubation with the substrate at room temperature.

Where referred to in the figures, the specific antibody titer is expressed as the reciprocal of the dilution showing activity above background levels.

EXAMPLE 4

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Bacterial Colonization

Bacteria were enumerated by spreading serial dilutions (10° to 10⁻³) directly onto TSA sheep blood agar plates followed by overnight incubation at 37°C, 5% CO₂. Colonization is defined as >500 cfu/ml of the challenge organism recovered.

To confirm that the bacteria recovered from milk secretions were *S. dysgalactiae*, selected colonies recovered from each animal were tested using an API strep-20 test (bioMerieux SA, Hazelwood, Missouri) according to the manufacturer's instructions. This test is a standardized method combining 20 biochemical tests for the determination of enzymatic activity and fermentation of sugars. The reactions are read according to a reading table and the identification is obtained by either referring to the analytical profile index or using identification software.

Following challenge, animals from all groups were shown to be colonized by S. dysgalactiae (Figure 20). Only the GapC-immunized cows had a statistically significant reduction in the number of infected quarters and total numbers of bacteria isolated per quarter. Therefore, immunization with GapC reduced bacterial colonization following challenge with S. dysgalactiae.

The relationship between anti-GapC titer and bacterial colonization is shown in Figures 21 and 22. There was a strong correlation between anti-GapC serum antibody level and the maximum number of bacteria (expressed in CFU (log₁₀)/ml milk) found in any



quarter r = 0.74) (Figure 21) as well as the total number of infected quarters ($r^2 = 0.74$) (Figure 22). Correlation was calculated using GraphPad Prism software, version 2.01 (GraphPad Software Inc., San Diego, California).

This correlation is also illustrated in Figure 23 and 24 where the GapC-immunized group is subdivided into high titer and low titer responders. In these figures, "low titer responders" refer to the four animals with the poorest response against GapC while "high titer responders" refer to the remainder of the group. No colonization occurred in the high titer group, while even the low titer group showed reduced numbers of bacteria recovered after day 3.

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EXAMPLE 5

Determination of inflammatory response

Inflammatory response was measured as a function of somatic cell count (i.e., lymphocytes, neutrophils, and monocytes). Somatic cell counts were measured in a Coulter counter using standard techniques, as recommended by Agriculture and Agri-Food Canada Pamphlet IDF50B (1985) *Milk and Milk products—Methods of Sampling*. Samples were always read within 48 hours of collection and fixation, at days 1 through 7 post challenge.

The numbers of somatic cells present in the gland was determined on each day post challenge. Numbers from the unchallenged quarter remained constant throughout the trial while on day 1, the GapC group was lower than the placebo-immunized group (Figure 25). The difference between the GapC and the placebo groups was statistically significant. The individual data from day 1 is shown in Figure 26; data for GapC-treated animals over a 7 day period post-challenge is shown in Figure 27. Samples from the quarters of GapC-immunized animals were indistinguishable from unchallenged quarters.

Therefore, immunization with GapC reduced the inflammatory response following challenge with S. dysgalactiae.

EXAMPLE 6

Cross-protection against S. uberis infection

During the last 3 days of the vaccine trial, a mixed population of bacteria were recovered from the mammary gland secretions. Further analysis with the API 20 Strep test confirmed the identity of the strains present in the mixture as S. dysgalactiae and S. uberis, the latter representing a natural infection which occurred during the trial. The group which

was immunized with GapC appeared to be significantly cross-protected against *S. uberis* colonization (see Figure 20), indicating that GapC may be a broadly cross-protective antigen capable of protecting against infection by multiple Streptococcal species.

Thus, vaccination with S. dysgalactiae GapC is therefore capable of providing cross-protection against S. uberis infection.

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Thus, the cloning, expression and characterization of various GapC plasmin binding proteins is disclosed, as are methods of using the same. Although preferred embodiments of the subject invention have been described in some detail, it is understood that obvious variations can be made without departing from the spirit and the scope of the invention as defined by the appended claims.

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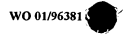
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What is claimed is:

- 1. An isolated GapC protein selected from the group consisting of:
- (a) an isolated *Streptococcus dysgalactiae* GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figure 1A-1B (SEQ ID NO:4);
- (b) an isolated Streptococcus agalactiae GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 2A-2B (SEQ ID NO:6);
- (c) an isolated *Streptococcus uberis* GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 3A-3B (SEQ ID NO:8);
- (d) an isolated *Streptococcus parauberis* GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 4A-4B (SEQ ID NO:10);
- (e) an isolated *Streptococcus iniae* GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 5A-5B (SEQ ID NO:12); and
- (f) immunogenic fragments of (a), (b), (c), (d) and (e) comprising at least about 5 amino acids.
 - 2. The isolated GapC protein of claim 1, wherein the protein is a *Streptococcus dysgalactiae* GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 1A-1B (SEQ ID NO:4).
 - 3. The isolated GapC protein of claim 1, wherein the protein is a *Streptococcus* agalactiae GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 2A-2B (SEQ ID NO:6).
- 4. The isolated GapC protein of claim 1, wherein the protein is a *Streptococcus* uberis GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 3A-3B (SEQ ID NO:8).



- 5. The isolated GapC protein of claim 1, wherein the protein is a *Streptococcus* parauberis GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 4A-4B (SEQ ID NO:10).
- 6. The isolated GapC protein of claim 1, wherein the protein is a *Streptococcus* iniae GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 5A-5B (SEQ ID NO:12).
- 7. An isolated polynucleotide comprising a coding sequence for an isolated GapC
 protein according to any of claims 1-6, or complements thereof.
 - 8. A recombinant vector comprising:
 - (a) the isolated polynucleotide of claim 7; and
- (b) at least one heterologous control element operably linked to said isolated
 polynucleotide, whereby said coding sequence can be transcribed and translated in a host cell, and at least one of said control elements is heterologous to said coding sequence.
 - 9. A host cell comprising the recombinant vector of claim 8.
- 20 10. A method for producing a GapC protein, said method comprising culturing the cells of claim 9 under conditions for producing said protein.
 - 11. A vaccine composition comprising a pharmaceutically acceptable vehicle and a GapC protein, wherein said GapC protein is selected from the group consisting of:
 - (a) a Streptococcus dysgalactiae GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 1A-1B (SEQ ID NO:4);
 - (b) a Streptococcus agalactiae GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 2A-2B (SEQ ID NO:6);
 - (c) a Streptococcus uberis GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 3A-3B (SEQ ID NO:8);
 - (d) a Streptococcus parauberis GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 4A-4B (SEQ ID NO:10);



- (e) a Streptococcus iniae GapC protein comprising the amino acid sequence shown at amino acid positions 1 to 336, inclusive, of Figures 5A-5B (SEQ ID NO:12);
- (f) a Streptococcus GapC protein having at least about 70% sequence identity to (a), (b), (c), (d) and (e); and
- (g) immunogenic fragments of (a), (b), (c), (d), (e) and (f), said fragments comprising at least about 5 amino acids.
- 12. The vaccine composition of claim 11, wherein said GapC protein comprises the amino acid sequence of the *Streptococcus dysgalactiae* GapC protein depicted at amino acid positions 1 to 336, inclusive, of Figures 1A-1B (SEQ ID NO:4) or an immunogenic fragment thereof, said fragment comprising at least about 5 amino acids.
- 13. The vaccine composition of claim 12, wherein said GapC protein comprises the amino acid sequence depicted at amino acid position 1 to 336, inclusive, of Figures 1A-1B (SEQ ID NO:4).
- 14. The vaccine composition of claim 11, wherein said GapC protein comprises the amino acid sequence of the *Streptococcus agalactiae* GapC protein depicted at amino acid positions 1 to 336, inclusive, of Figures 2A-2B (SEQ ID NO:6) or an immunogenic fragment thereof, said fragment comprising at least about 5 amino acids.
- 15. The vaccine composition of claim 14, wherein said GapC protein comprises the amino acid sequence depicted at amino acid positions 1 to 336, inclusive, of Figures 2A-2B (SEQ ID NO:6).
- 16. The vaccine composition of claim 11, wherein said GapC protein comprises the amino acid sequence of the *Streptococcus uberis* GapC protein depicted at amino acid positions 1 to 336, inclusive, of Figures 3A-3B (SEQ ID NO:8) or an immunogenic fragment thereof, said fragment comprising at least about 5 amino acids.
- 17. The vaccine composition of claim 16, wherein said GapC protein comprises the amino acid sequence depicted at amino acid positions 1 to 336, inclusive, of Figures 3A-3B (SEQ ID NO:8).

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18. The vaccine composition of claim 11, wherein said GapC protein comprises the amino acid sequence of the *Streptococcus parauberis* GapC protein depicted at amino acid positions 1 to 336, inclusive, of Figures 4A-4B (SEQ ID NO:10) or an immunogenic fragment thereof, said fragment comprising at least about 5 amino acids.

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19. The vaccine composition of claim 18, wherein said GapC protein comprises the amino acid sequence depicted at amino acid positions 1 to 336, inclusive, of Figures 4A-4B (SEQ ID NO:10).

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20. The vaccine composition of claim 11, wherein said GapC protein comprises the amino acid sequence of the *Streptococcus iniae* GapC protein depicted at amino acid positions 1 to 336, inclusive, of Figures 5A-5B (SEQ ID NO:12) or an immunogenic fragment thereof, said fragment comprising at least about 5 amino acids.

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21. The vaccine composition of claim 20, wherein said GapC protein comprises the amino acid sequence depicted at amino acid positions 1 to 336, inclusive, of Figures 5A-5B (SEQ ID NO:12).

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- 22. The vaccine composition according to any of claims 11-21, further comprising an adjuvant.
 - 23. A method of producing a vaccine composition comprising the steps of
- (1) providing a GapC protein or an immunogenic fragment thereof, said fragment comprising at least about 5 amino acids, and

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- (2) combining said protein with a pharmaceutically acceptable vehicle.
- 24. The method of claim 23, wherein the GapC protein is the protein of any of claims 1-6.

- 25. Antibodies directed against the isolated GapC protein of any of claims 1-6.
- 26. The antibodies of claim 25 wherein said antibodies are polyclonal.

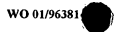
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- 27. The antibodies of claim 25 wherein said antibodies are monoclonal.
- 28. Use of a GapC protein according to any of claims 1-6, in the manufacture of a vaccine composition useful for treating or preventing a bacterial infection in a vertebrate subject.
- 29. The use of claim 28, wherein said bacterial infection is a streptococcus infection.
 - 30. The use of claim 29, wherein said bacterial infection causes mastitis.
- 31. Use of a polynucleotide according to claim 7 in the manufacture of a vaccine composition for treating or preventing a bacterial infection in a vertebrate subject.
 - 32. The use of claim 31, wherein said bacterial infection is a streptococcal infection.
 - 33. The use of claim 32, wherein said bacterial infection causes mastitis.
- 34. Use of a GapC protein in the manufacture of a diagnostic reagent for detecting Streptococcus antibodies in a biological sample.
 - 35. Use of a GapC protein according to any of claims 1-6 in the manufacture of a diagnostic reagent for detecting *Streptococcus* antibodies in a biological sample.
 - 36. Use of antibodies according to any of claims 25-27 in the manufacture of a diagnostic reagent for detecting a GapC protein in a biological sample.
 - 37. An immunodiagnostic test kit for detecting *Streptococcus* infection, said test kit comprising a GapC protein and instructions for conducting the immunodiagnostic test.
 - 38. An immunodiagnostic test kit for detecting *Streptococcus* infection, said test kit comprising antibodies directed against a GapC protein and instructions for conducting the immunodiagnostic test.



- 39. A method of treating or preventing a bacterial infection in a vertebrate subject comprising administering to said subject a therapeutically effective amount of a vaccine composition according to any of claims 11-22.
- 5 40. The method of claim 39, wherein said bacterial infection is a streptococcus infection.
 - 41. The method of claim 40, wherein said bacterial infection causes mastitis.
- 42. A method of treating or preventing a bacterial infection in a vertebrate subject comprising administering to said subject a therapeutically effective amount of a polynucleotide according to claim 7.
- 43. The method of claim 42, wherein said bacterial infection is a streptococcal infection.
 - 44. The method of claim 43, wherein said bacterial infection causes mastitis.
- 45. A method of detecting *Streptococcus* antibodies in a biological sample, comprising:
 - (a) reacting said biological sample with an isolated GapC protein under conditions which allow said *Streptococcus* antibodies, when present in the biological sample, to bind to said GapC protein to form an antibody/antigen complex; and
 - (b) detecting the presence or absence of said complex, thereby detecting the presence or absence of *Streptococcus* antibodies in said sample.
 - 46. A method of detecting *Streptococcus* antibodies in a biological sample, comprising:
- (a) reacting said biological sample with an isolated GapC protein according to any of claims 1-6, under conditions which allow said Streptococcus antibodies, when present in the biological sample, to bind to said GapC protein to form an antibody/antigen complex; and
 - (b) detecting the presence or absence of said complex, thereby detecting the presence or absence of *Streptococcus* antibodies in said sample.



- 47. A method of detecting a GapC protein in a biological sample, comprising:
- (a) reacting said sample with antibodies directed against the GapC protein under conditions which allow said antibodies to bind to said GapC protein, when present in the sample, to form an antibody/antigen complex; and
- (b) detecting the presence or absence of said complex, thereby detecting the presence or absence of a GapC protein in said sample.

atg Met 1	gta Val	gtt Val	aaa Lys	gtt Val 5	ggt Gly	att Ile	aac Asn	ggt Gly	ttc Phe 10	ggt Gly	cgt Arg	atc Ile	gga Gly	egt Arg 15	ctt Leu	48
gca Ala	ttc Phe	cgt Arg	cgt Arg 20	att Ile	caa Gln	aat Asn	gtt Val	gaa Glu 25	ggt Gly	gtt Val	gaa Glu	gta Val	act Thr 30	cgt Arg	atc Ile	96
aac Asn	gac Asp	ctt Leu 35	aca Thr	gat Asp	cca Pro	aac Asn	atg Met 40	ctt Leu	gca Ala	cac His	ttg Leu	ttg Leu 45	aaa Lys	tac Tyr	gat Asp	144
aca Thr	act Thr 50	caa Gln	gga Gly	cgt Arg	ttt Phe	gac Asp 55	gga Gly	act Thr	gtt Val	gaa Glu	gtt Val 60	aaa Lys	gaa Glu	ggt	gga Gly	192
ttt Phe 65	gaa Glu	gta Val	aac Asn	gga Gly	aac Asn 70	ttc Phe	atc Ile	aaa Lys	gtt Val	tct Ser 75	gct Ala	gaa Glu	cgt Arg	gat Asp	cca Pro 80	240
gaa Glu	aac Asn	atc Ile	gac Asp	tgg Trp 85	gca Ala	act Thr	gac Asp	ggt Gly	gtt Val 90	gaa Glu	atc Ile	gtt Val	ctg Leu	gaa Glu 95	gca Ala	288
act Thr	ggt Gly	ttc Phe	ttt Phe 100	gct Ala	aaa Lys	aaa Lys	gaa Glu	gct Ala 105	gct Ala	gaa Glu	aaa Lys	cac His	tta Leu 110	cat His	gct Ala	336
aac Asn	ggt Gly	gct Ala 115	aaa Lys	aaa Lys	gtt Val	gtt Val	atc Ile 120	aca Thr	gct Ala	cct Pro	ggt Gly	gga Gly 125	aac Asn	gac Asp	gtt Val	384
											ctt Leu 140					432
aca Thr 145	Val	atc Ile	tca Ser	ggt Gly	gct Ala 150	tca Ser	tgt Cys	act Thr	aca Thr	aac Asn 155	tgt Cys	tta Leu	gct Ala	cct Pro	atg Met 160	480
gct Ala	aaa Lys	gct Ala	ctt Leu	cac His 165	Asp	gca Ala	ttt Phe	ggt Gly	ato Ile 170	Gli	aaa Lys	ggt Gly	ctt Leu	atg Met 175	Thr	528
aca Thr	atc Ile	cac His	gct Ala 180	Tyr	act Thr	ggt	gac Asp	caa Gln 185	Met	ato : Ile	ctt Leu	gac Asp	gga Gly 190	Pro	cac His	576
cgt Arg	ggt Gly	ggt Gly 199	Asp	ctt Leu	cgt Arg	cgt Arg	gct Ala 200	Arg	gct Ala	ggt a Gly	gct Ala	gca Ala 205	Asr	att Ile	gtt Val	624

FIG. 1A



		Ser														672
ttg Leu 225	aat Asn	ggt Gly	aaa Lys	ctt Leu	gat Asp 230	ggt Gly	gct Ala	gca Ala	caa Gln	cgt Arg 235	gtt Val	cct Pro	gtt Val	cca Pro	act Thr 240	720
gga Gly	tca Ser	gta Val	act Thr	gag Glu 245	ttg Leu	gtt Val	gta Val	act Thr	ctt Leu 250	gat Asp	aaa Lys	aac Asn	gtt Val	tct Ser 255	gtt Val	768
gac Asp	gaa Glu	atc Ile	aac Asn 260	gct Ala	gct Ala	atg Met	aaa Lys	gct Ala 265	gct Ala	tca Ser	aac Asn	gac Asp	agt Ser 270	ttc Phe	ggt Gly	816
tac Tyr	act Thr	gaa Glu 275	gat Asp	cca Pro	att Ile	gtt Val	tct Ser 280	tca Ser	gat Asp	atc Ile	gta Val	ggc Gly 285	gtg Val	tca Ser	tac Tyr	864
ggt Gly	tca Ser 290	ttg Leu	ttt Phe	gac Asp	gca Ala	act Thr 295	Gln	act Thr	aaa Lys	gtt Val	atg Met 300	gaa Glu	gtt Val	gac Asp	gga Gly	912
tca Ser 305	Gln	ttg Leu	gtt Val	aaa Lys	gtt Val 310	gta Val	tca Ser	tgg Trp	tat Tyr	gac Asp 315	Asn	gaa Glu	atg Met	tct Ser	tac Tyr 320	960
act Thr	gct Ala	caa Gln	'ctt Leu	gtt Val 325	Arg	aca Thr	ctt Leu	gag Glu	tac Tyr 330	ttt Phe	gca Ala	aaa Lys	atc Ile	gct Ala 335	aaa Lys	1008
taa																1011

FIG. 1B

atg Met 1	gta Val	gtt Val	aaa Lys	gtt Val 5	ggt Gly	att Ile	aac Asn	ggt Gly	ttc Phe 10	ggt Gly	cgt Arg	atc Ile	ggt Gly	cgt Arg 15	ctt Leu	48
gca Ala	ttc Phe	cgt Arg	cgc Arg 20	atc Ile	caa Gln	aac Asn	gta Val	gaa Glu 25	ggt Gly	gtt Val	gaa Glu	gtt Val	act Thr 30	cgt Arg	atc Ile	96
aac Asn	gac Asp	ctt Leu 35	aca Thr	gat Asp	cca Pro	aac Asn	atg Met 40	ctt Leu	gca Ala	cac His	ttg Leu	ttg Leu 45	aaa Lys	tat Tyr	gac Asp	144
aca Thr	act Thr 50	caa Gln	ggt Gly	cgt Arg	ttc Phe	gac Asp 55	ggt Gly	act Thr	gtt Val	gaa Glu	gtt Val 60	aaa Lys	gaa Glu	ggt Gly	gga Gly	192
ttc Phe 65	gaa Glu	gtt Val	aac Asn	ggt Gly	caa Gln 70	ttt Phe	gtt Val	aaa Lys	gtt Val	tct Ser 75	gct Ala	gaa Glu	ege Arg	gaa Glu	cca Pro 80	240
gca Ala	aac Asn	att Ile	gac Asp	tgg Trp 85	gct Ala	act Thr	gat Asp	ggc Gly	gta Val 90	gaa Glu	atc Ile	gtt Val	ctt Leu	gaa Glu 95	gca Ala	288
act Thr	ggt Gly	ttc Phe	ttt Phe 100	gca Ala	tca Ser	aaa Lys	gaa Glu	aaa Lys 105	gct Ala	gga Gly	caa Gln	cac His	atc Ile 110	cat His	gaa Glu	336
aat Asn	ggt Gly	gct Ala 115	aaa Lys	aaa Lys	gtt Val	gtt Val	atc Ile 120	aca Thr	gct Ala	cct Pro	ggt Gly	gga Gly 125	aac Asn	gac Asp	gtt Val	384
aaa Lys	aca Thr 130	gtt Val	gtt Val	ttc Phe	aac Asn	act Thr 135	aac Asn	cac His	gat Asp	atc Ile	ctt Leu 140	gat Asp	gga Gly	act Thr	gaa Glu	432
aca Thr 145	gtt Val	atc Ile	tca Ser	ggt Gly	gct Ala 150	tca Ser	tgt Cys	act Thr	aca Thr	aac Asn 155	tgt Cys	ctt Leu	gct Ala	cca Pro	atg Met 160	480
gct Ala	aaa Lys	gct Ala	tta Leu	caa Gln 165	gac Asp	aac Asn	ttt Phe	ggt Gly	gtt Val 170	Lys	caa Gln	ggt Gly	ttg Leu	atg Met 175	act Thr	528
act Thr	atc :Ile	cac His	gca Ala 180	Tyr	act Thr	ggt	gac gac	caa Gln 185	Met	atc Ile	ctt Leu	gac Asp	gga Gly 190	cca Pro	cac His	576
cgt Arg	ggt	ggt Gly 195	Asp	ctt Leu	cgt Arg	cgt Arg	gct Ala 200	Arg	gca Ala	ggt Gly	gct Ala	gca Ala 205	aac Asn	atc Ile	gtt Val	624

FIG. 2A



PLO	210	ser	Thr	GIĄ	gct Ala	Ala 215	ГÀВ	Ala	Ile	Gly	Leu 220	Val	Ile	Pro	Glu	672
ttg Leu 225	aac Asn	ggt Gly	aaa Lys	ctt Leu	gat Asp 230	ggt Gly	gct Ala	gca Ala	caa Gln	cgt Arg 235	gtt Val	cct Pro	gtt Val	cca Pro	act Thr 240	720
gga Gly	tca Ser	gta Val	act Thr	gaa Glu 245	ttg Leu	gtt Val	gca Ala	act Thr	ctt Leu 250	gaa Glu	aaa Lys	gac Asp	gta Val	act Thr 255	gtc Val	768
gaa Glu	gaa Glu	gta Val	aat Asn 260	gca Ala	gct Ala	atg Met	aaa Lys	gca Ala 265	gca Ala	gct Ala	aac Asn	gat Asp	tca Ser 270	tac Tyr	Gly Gly	816
tat Tyr	act Thr	gaa Glu 275	gat Asp	cca Pro	atc Ile	gta Val	tca Ser 280	tct Ser	gat Asp	atc Ile	gtt Val	ggt Gly 285	att Ile	tca Ser	tac Tyr	864
ggt Gly	tca Ser 290	ttg Leu	ttt Phe	gat Asp	gct Ala	act Thr 295	caa Gln	act Thr	aaa Lys	gtt Val	caa Gln 300	act Thr	gtt Val	gac Asp	ggt Gly	912
aac Asn 305	caa Gln	ttg Leu	gtt Val	aaa Lys	gtt Val 310	gtt Val	tca Ser	tgg Trp	tac Tyr	gat Asp 315	aac Asn	gaa Glu	atg Met	tca Ser	tac Tyr 320	960
act Thr	tca Ser	caa Gln	ctt Leu	gtt Val 325	cgt Arg	aca Thr	ctt Leu	gag Glu	tac Tyr 330	ttt Phe	gca Ala	aaa Lys	atc Ile	gct Ala 335	aaa Lys	1008
taa																1011

FIG. 2B

															-	
atg Met 1	gta Val	gtt Val	aaa Lys	gtt Val 5	ggt	att Ile	aac Asn	ggt Gly	ttc Phe 10	ggt Gly	cgt Arg	atc Ile	gga Gly	cgt Arg 15	ctt Leu	48
				att Ile												96
aac Asn	gat Asp	ctt Leu 35	act Thr	gac Asp	cca Pro	aat Asn	atg Met 40	ctt Leu	gca Ala	cac His	ttg Leu	ttg Leu 45	aaa Lys	tat Tyr	gat Asp	144
				cgt Arg												192
				gga Gly												240
				tgg Trp 85												288
				gct Ala												336
			Lys	aaa Lys				Thr								384
		Val		ttt. Phe			Asn					Asp				432
	. Val			ggt Gly		Ser					Cys					480
gct Ala	aaa Lys	gct Ala	tte Lei	g caa ı Glr 165	l Asi	aac Asn	ttt Phe	ggt Gly	gti Y Val	l Ly	a caa s Gli	a ggt a Gly	ttg Lev	atg Met 175	g aca : Thr	528
act Thi	t ato	cac His	get 8 Ala 180	а Туз	c act	ggt Gly	ga (: Asj	c cas o Gli 18!	n Me	g ato	c cti e Lei	t gad ı Asp	99a Gly	Pro	a cac o His	576
cg:	t ggi g Gly	ggt y Gl; 19:	y As	c ctt p Lei	ı Arg	g Arg	g Ala 20	a Ar	t gc g Ala	t gg a Gl	t gca y Ala	a ago a Sei 209	: Ası	c ati	t gtt e Val	624

FIG. 3A



Pro		tca Ser													672
		ggt Gly			_		_	_		_	_		_		720
		gta Val													768
_	_	atc Ile		_	_	_		_	_	_		_			 816
		gaa Glu 275	_			_			_				_	_	864
		ttg Leu		-	_					_			-	_	 912
	Gln	tta Leu	_		_	_				_	Asn	_	_		960
	_	caa Gln		_	Arg					Phe	_			_	1008
taa															1011

FIG. 3B



atg Met 1	gta Val	gtt Val	Lys	gtt Val 5	ggt Gly	att Ile	aac Asn	ggt Gly	ttt Phe 10	ggc Gly	cgt Arg	atc Ile	gga Gly	cgt Arg 15	ctt Leu	48
gct Ala	ttc Phe	cgt Arg	cgt Arg 20	att Ile	caa Gln	aat Asn	gta Val	gaa Glu 25	ggt Gly	gtt Val	gaa Glu	gtt Val	act Thr 30	cgc Arg	atc Ile	96
aac Asn	Asp Asp	ctt Leu 35	aca Thr	gat Asp	cca Pro	aat Asn	atg Met 40	ctt Leu	gca Ala	cac His	ttg Leu	tta Leu 45	aaa Lys	tac Tyr	gat Asp	144
								act Thr								192
								aaa Lys								240
								ggt Gly								288
				Ala				gct Ala 105								336
			Lys					Thr							gtg Val	384
		Val					Asn					Asp			gaa Glu	432
	Val					Ser					Cys				atg Met 160	480
			. Let		ı Ası					Lys					act Thr	528
				а Туз					ı Met					Pro	cac His	576
			/ Asj					a Arg					a Ası		t gtt e Val	624

FIG. 4A



Pro							gca Ala							672
				-		_	gca Ala		_	-		_		720
							gtt Val							768
_	_			_			gct Ala 265	-	_		_			816
		_	_		_		tct Ser	~		-		_		864
							act Thr							912
	Gln		_	_	_		tgg Trp		_		_	_		960
				Arg			gag Glu							1008
taa	•													1011

FIG. 4B

_	gta Val	-		_			•				_			_		48
	ttc Phe															96
	gac Asp															144
	act Thr 50															192
	gaa Glu															240
gca Ala	aac Asn	att Ile	gac Asp	tgg Trp 85	gct Ala	act Thr	gat Asp	ggt Gly	gta Val 90	Asp	atc Ile	gtt Val	ctt Leu	gaa Glu 95	gca Ala	288
	ggt Gly			Ala					Ala					His		336
	e ggt n Gly		Lys					Thr					Asn			384
aaa Ly:	a aca Thi	Val	gtt L Val	tac L Tyr	aac Asr	act Thr 135	Asn	cat His	gat Asr	att Ile	ctt Lev 140	ı Asp	gga Gly	act Thr	gaa Glu	432
	r Va					s Ser					в Су				atg Met 160	480
					n Āsī					l Ly					g act t Thr	528
				у Ту					n Me					y Pro	a cac o His	576
			y As					a Ar					a As		c gtt e Val	624

FIG. 5A



cct Pro	aac Asn 210	tca Ser	act Thr	ggt Gly	gct Ala	gct Ala 215	aaa Lys	gca Ala	atc Ile	ggt Gly	ctt Leu 220	gtt Val	atc Ile	cca Pro	gaa Glu	672
tta Leu 225	aat Asn	ggt Gly	aaa Lys	ctt Leu	gac Asp 230	ggt Gly	gct Ala	gca Ala	caa Gln	cgt Arg 235	gtt Val	cct Pro	gtt Val	cca Pro	act Thr 240	720
gga Gly	tca Ser	gta Val	act Thr	gaa Glu 245	tta Leu	gta Val	gca Ala	gtt Val	ctt Leu 250	gaa Glu	aaa Lys	gat Asp	act Thr	tca Ser 255	gta Val	768
gaa Glu	gaa Glu	atc Ile	aat Asn 260	gca Ala	gct Ala	atg Met	ааа Ьув	gca Ala 265	gca Ala	gct Ala	aac Asn	gat Asp	tca Ser 270	tac Tyr	ggt Gly	816
tac Tyr	act Thr	gaa Glu 275	Asp	gct Ala	atc Ile	gta Val	tca Ser 280	Ser	gat Asp	atc Ile	gta Val	ggt Gly 285	Ile	tct Ser	tac	864
ggt Gly	tca Ser 290	Lev	ttt Phe	gat Asp	gct Ala	act Thr 295	Gln	act Thr	aaa Lys	gta Val	caa Glm 300	Thr	gtt Val	gat	gga	912
aat Asi 30!	ı Glr	ı tt <u>e</u> ı Leı	g gtt n Val	aaa Lys	gtt Val	. Val	tca Ser	tgg Trp	tat Tyr	gac Asp 315) Asi	gaa Glu	atg Met	tct Ser	tac Tyr 320	960
act Th	t gct r Ala	caa a Gl	a cti n Lei	t gtt 1 Va: 32!	l Arg	act Thi	c ctt	gag 1 Glu	tac Tyi	Phe	gca Ala	a aaa a Lys	a ato	gct Ala 339	aaa Lys	1008 [.]
ta	a															1011

FIG. 5B

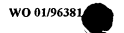
	1				50
DysGapC	ATGGTAGTTA	AAGTTGGTAT	TAACGGTTTC	GGTCGTATCG	GACGTCTTGC
SpyGapC					
SeqGapC					
			t		
AgalGapCDNA					
BovGapC		~~~~~~		~~~c	-gcg-t
	51				100
			TTGAAGGTGT		
SpyGapC	c	ca	-c		t-
ParaUbGapC	t		-a	t	C
UberGapc		c-			t
AgalGapCDNA	C	CC-	-a	t	
SiniGapC					t-
BovGapC	cac-a-ggc-	gc-tttt	cgcaaa	gca-cgtc	gcct-
			_		
	101				150
DysGapC	ACCTTAC	AGATCCAAAC	ATGCTTGCAC	ACTTGTTGAA	ATACGATACA
SpyGapC		t			c
SeqGapC	,		-		c
ParaUbGapC		t		a	c
UberGapc	-t	tct			t
AgalGapCDNA					tc
SiniGapC		t			t
RoyGanC	cct-c-t	ta-tta	. taga_g_tgt	2	

FIG. 6A



	151				222
DecCanC			2 2 CONCOUNT	G0000000000000000000000000000000000000	200
			AACTGTTGAA		
			aa		
			at		
			ta		
UberGapc		c	a	t-	C
AgalGapCDNA		c	t		C
SiniGapC			a	t-	C
BovGapC	cca	agca	caca-g	-cag-ga-c-	-gaagc-c-t
	201				250
			AAGTTTCTGC		
SpyGapC	a			t	
SeqGapC	a			t	
ParaUbGapC	c	at-		aaa	c-a-
UberGapc				aaa	
AgalGapCDNA					
SiniGapC		-qtq-t-		aca	c
			cca-c-tcca		
-		33			
	251				300
DysGapC	TCGACTGGGC	AACTGACGGT	GTTGAAATCG	TTCTGGAAGC	AACTGGTTTC
			a		
AgalGapCDNA					
			ac		
bovdapc	-ca-gg	tgact	-cgtat-	-ag-ggc-	cgg
	301				250
Dreadand		** C * * C C C C C C C C C C C C C C C	maxxxx	mms as maams	350
			TGAAAAACAC		
			t		
			t		
AgalGapCDNA	atc	aaa	g-c	a-caa-	-t
			c		
BovGapC	cacc-	tggaag	gggct	ga-g-g	cc
	351				400
			CTGGTGGAAA		
			g-		
			g-		
BovGap	C g-ggca-c	:t-ta-	tc	:t-cccc	: -tgtga

FIG. 6B



	401				
DvsGapC		CCVCCV, CV	TCTTGACGGT	3.00000000	450
SpyGapC		CCACGA.CAT	TCTTGACGGT	ACIGAAACAG	TTATCTCAGG
SegGapC					
ParaUbGapC	-t	t- +			
UberGanc	-t		cta		t
AgalGapCDNA	7777			at-	-at
AgalGapCDNA SiniGapC	3		cta		
BoyGanC	-ccc-ctc-		ta		
Dovempe	-999-909	a-g	a-aaac	cctc-aga	gagcaa
				•	
	451				
DysGapC	•	A CTACK A A COL	Commit domos	M1 M00000000000000000000000000000000000	500
SpyGapC		ACIACAMACI	GTTTAGCTCC	TATGGCTAAA	GCTCTTCACG
SecGapC				£	c-tc-
ParaUbGapC				C	c-tc-
UberGano					a
AgalGanCDNA					g
AgalGapCDNA SiniGanC			c-t		a
PovCapC		_			aa
bovdapc	66	CC	-cgc	ccg	-tca-ct-
	501				
DvsGapC		ሞሽጥ////ሽሽሽሽ	Component man		550
SnyGanC	AIGCAIIIGG	IAICCAAAAA	GGTCTTATGA	CTACAATCCA	CGCTTATACT
ექანება	902	-aca	c	a	
Parattheane	902	-aca	c	a	
TherCane		cg-a	t-a	a	
Japleson		-g	t-g	-at	
arrechemus	-C	-g	t-g	t	a
DIMIGAPC		-g-a	t-a	t	t-g
BovGapC	-cc	ca-cgtgg-g	ac	-ctg	cat
	551				
DucGanc		MG3 manner			600
೨ರ್ಥವಿಗಳ	GGIGACCAAA	TGATCCTTGA	CGGACCACAC	CGTGGTGGTG	ACCTTCGTCG
SpyGapC					
Sequapo		g	tac-gt	g	-t
Far anngahe	E	C-t	tt		+
UberGapc AgalGapCDNA					
AgalGapCDNA					
STHIGAPC		g-t			-t
BovGapC	-ccacg-	actg-g	tcctc-	gaagc	tgtggga
					- 35
	601 .				
Decodora		00000			650
Dy scapt	recreatect	GGIGCIGCAA	ACATTGTTCC	TAACTCAACT	GGTGCTGCTA
So-Co-C	aC				
Darathcase					cg
The work = -	6	aac-	-tt		
yda j Gardina onergabc		aagc-	t		
**Aaraabcn##	a	~			
STHIGAPC	a	-ca	~		
DOVGADC	c-gca-gg	-cccag-	-ta-c	-gctt	CC-

FIG. 6C



	651				
DvsGapC		יאיי פינייניבארייריאיי	CCAGAATTGA	ስጥር-ርጥአ አ አ ረ ጥ	700
SpyGapC		TOTIGITATE	c-t-	AIGGIAAACI	IGATGGTGCT
SegGapC			g-	-C	
ParaUbGapC	a		ta-	-t	
UberGapc	a		a-	-t	
AgalGapCDNA		a		-C	
SiniGapC	a		a-	-t	
BovGapC	-qcq-q	caagc	tgc-c-	-CGG	cactcate
-		-	- 500	• 99	cactcatg
	701				750
DysGapC	GCACAACGTG	TTCCTGTTCC	AACTGGATCA	GTAACTGAGT	TGGTTGTAAC
SpyGapC				a-	
SeqGapC				a-	t
ParaUbGapC		-aa	at	a	-aaat
UberGapc					-aaat
AgalGapCDNA	~				90
SiniGapC					-aaat
BovGapC	cttcc-	-ccac	c-ac-tgt	tatatc	acctacca
				-505 00	
	751				800
DysGapC	TCTTGATAAA	AACGTTTCTG	TTGACGAAAT	CAACGCTGCT	
SpyGapC	c	at	c	t-t	
SeqGapC	c	at	c	t	
ParaUbGapC	a-t	aaca-	-a	tta	
UberGapc		aaca-		a	
AgalGapCDNA		taa	-ca-	at	
SiniGapC		taca-	-a	t	
BovGapC	cgg	cct-ccaaqt	atg	gaag-tg	ggcag-
				35	y godg
	801				850
DysGapC	CTTCAAACGA	CAGTTTCGGT	TACACTGAAG	ATCCAATTGT	
SpyGapC	t	-agc-t			teeseesee
SegGapC	t	-agc-t			t
ParaUbGapC	-ag-tt	at	·		
UberGapc	q	aa		-C	at
AgalGapCDNA	-aq-t	a	t		
SiniGapC	-aq-t	a		a-t-i	
BovGapC	-ata-a-a	cc-tct-aag	ggt-ct	gctac-ct-a	anacraa-t-
~	, ,,,		35 0 00	30000 00 a	ggaccag-t-
	851				900
DysGapC	ATCGTAGGCG	TGTCATA	CGGTTCATTG	TTTGACGCAA	
SpyGapC	cg	-a		ca-	
ParaUbGapC	ta	t-t	a	C	
UberGapo	a-cta	a-t			
AgalGapCDNA	tta	-t			
SiniGapC	ta	-tt			
BovGapC	gteet-eg	ac-tca-cag	a-a-tcac	-c-tc-a-ct	tca-taaa

FIG. 6D

	901	-			950
DysGapC	AGTTATGGAA	GTTGACGGAT	CACAATTGGT	TAAAGTTGTA	TCATGGTATG
SpyGapC	aatggaa	t	ca	a	
				a	
ParaUbGapC					
AgalGapCDNA					
				cgc-ca	
_		_		_	
•					
	951				1000
DysGapC	ACAATGAAAT	GTCTTACACT	GCTCAACTTG	TTCGTACACT	TGAGTATTTT
				-at	
				a	
				t	
AgalGapCDNA					
SiniGapC	t			t	
		-33- 3-			
	1001	101	.8		
DvsGapC	GCAAAAATCG	СТАЛАТАЛ	-		
	t-				
	,				
AgalGapCDNA					
Barranc					

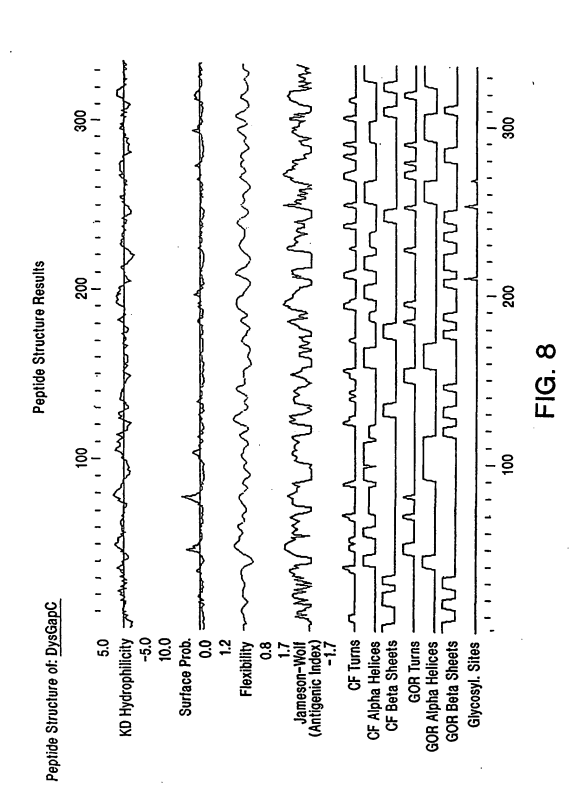
FIG. 6E

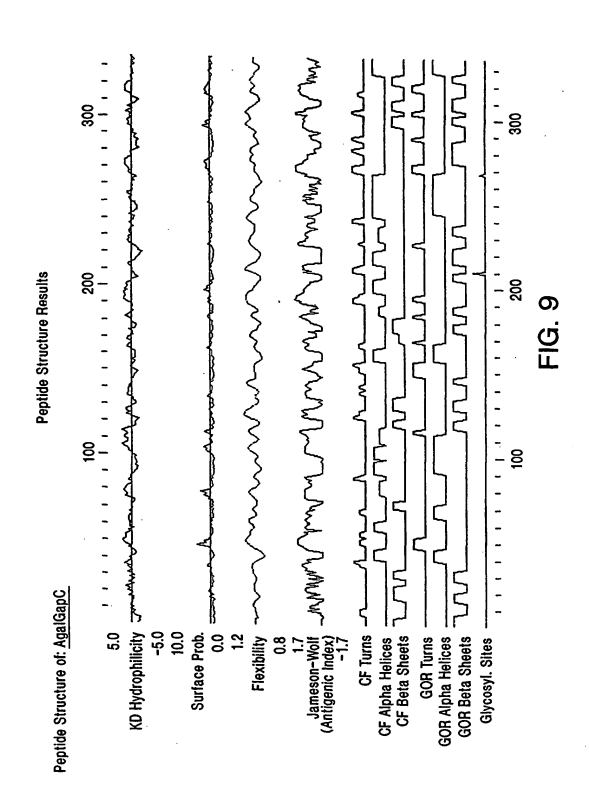
	1				50
DysGapC	MVVKVGINGF	GRIGRLAFRR	IQNVEGVEVT	RIND.LTDPN	MLAHLLKYDT
SpyGapC			I		
SeqGapC					
PUberGapC					
UberGapC	• • • • • • • • • • • • • • • • • • •				
AgalGapC					
IniaeGapC					
BovGapC		vt-a	af-sgk-div	apfi-lh	ymvymfqs
_		•			
	51				100
		VKEGGFEVNG	NFIKVSAERD	PENIDWATDG	VEIVLEATGF
EpyGapC					
SeqGapC					
PUberGapC		dd	kk-	q	
UberGapC		d	k-		
AgalGapC			q-ve	-a	-d
IniaeGapC		d	s-ve	-a	-d
BovGapC	-h-k-nk	aen-klvi	ka-tifq	-ak-gda-	a-y-v-sv
					150
	101				150
DysGapC	FAKKEAAEK	LHANGAKKVV	ITAPGGNDV	TAALMINEDI	LDGTETVISG
SpyGapC				-16-1144-	
SeqGapC	E)		directors-	
Pubergapo	a	e	a		
UperGapC	:a	·	·a		
AgalGapC	ek-gq	- 1-e			
rulaeGapC	:s-aq·	. 1		y	12-i
BovGapC	: -ccm-K-ga	кgr-:	ssaa <u>r</u>	mr-mgvex	ynn-lkiv-n

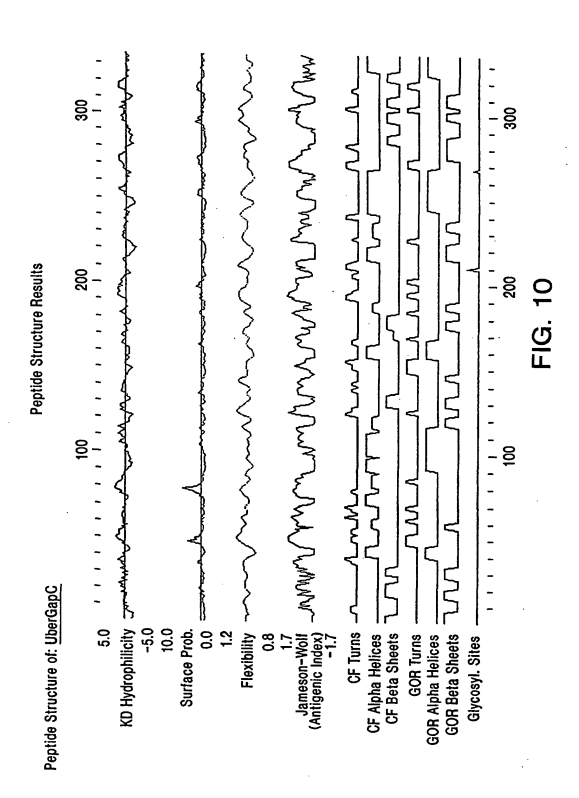
FIG. 7A

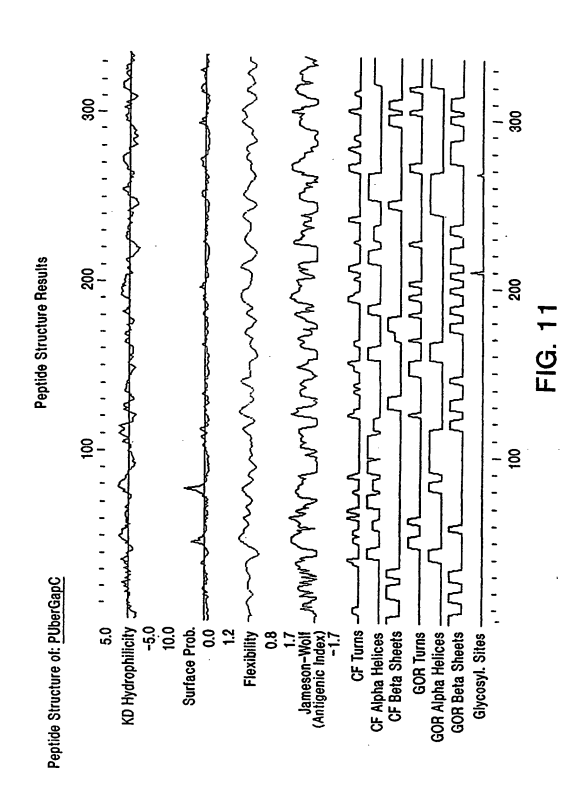
	151				200
DysGapC	ASCTINCLAP			AYTGDQMILD	GPHRGGDLRR
SeqGapC				v-	-hrg
PUberGapC		q-n	v		
UberGapC		q-n	v		
AgalGapC		q-n 	v		
IniaeGapC		q-n	V	gv	
BovGapC		1vih-h	ivev-	-i-at-ktv-	s-klw-d
	201				250
	ARAGAANIVP				
SpyGapC					
SeqGapC		r			
PUberGapC	n				av
UberGapC	8				av
AgalGapC					a-
IniaeGapC	a				av
BovGapC	g-ga-qi-	av-	k	t-m-ft-	nvvd-tcr
	^-7				
D	251				300
	LDKNVSVDEI				VSYGSLFDAT
SpyGapC	LDKNVSVDEI	-6			VSYGSLFDAT
SpyGapC SeqGapC	LDKNVSVDEI	-6			VSYGSLFDAT
SpyGapC SeqGapC PUberGapC	LDKNVSVDEI	-sva	y		VSYGSLFDAT
SpyGapC SeqGapC PUberGapC UberGapC	LDKNVSVDEI	a-	y	i-	VSYGSLFDAT
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC	LDKNVSVDEIn-etee-ete	-sva	 y y	1	VSYGSLFDAT m-f i
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC	LDKNVSVDEIn-etee-etee-d-te-v -e-dte	-6a -sva a	y y y	i-	VSYGSLFDAT m-f i
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC	LDKNVSVDEIn-etee-ete	-6a -sva a	y y y	i-	VSYGSLFDAT m-f i
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC	LDKNVSVDEIn-etee-etee-d-te-v -e-dte	-6a -sva a	y y y	i-	VSYGSLFDAT m-f i
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC	LDKNVSVDEIn-etee-etee-d-tee-paky	-6a -sva a	y y y	ia	w-f ma i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC	LDKNVSVDEI	-8 -sva a a kkvv-qeg	y y y y plkgilg	i i -a	VSYGSLFDAT m-f i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC	LDKNVSVDEI	-sa -sva a a kkvv-qeg	yyy plkgilg	-a	VSYGSLFDAT m-f ma i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC	LDKNVSVDEI	-sa -sva a a kkvv-qeg	yyy plkgilg	-a	VSYGSLFDAT m-f ma i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC DysGapC SpyGapC SeqGapC	LDKNVSVDEI	-8a -sva a a kkvv-qeg	y y y y plkgilg	-a	VSYGSLFDAT m-f ma i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC DysGapC SpyGapC SeqGapC	LDKNVSVDEI -n-etee-etee-dtee-paky 301 QTKVMEVDGS	-8a -sva a a kkvv-qeg	d	-a	VSYGSLFDAT m-f ma i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC DysGapC SpyGapC SeqGapC PUberGapC	LDKNVSVDEI -n-etee-etee-dtee-paky 301 QTKVMEVDGS	-8aa	plkgilg	-a	VSYGSLFDAT m-f ma i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC DysGapC SpyGapC SeqGapC PUberGapC AgalGapC	LDKNVSVDEI -n-etee-etee-dtee-paky 301 QTKVMEVDGSqtqtqt	-8aa	plkgilg	-qvc-fns	VSYGSLFDAT m-f i dths-tg
SpyGapC SeqGapC PUberGapC UberGapC AgalGapC IniaeGapC BovGapC SpyGapC SpyGapC SeqGapC PUberGapC UberGapC	LDKNVSVDEI -n-etee-etee-dtee-paky 301 QTKVMEVDGS	-8a	plkgilg	i -aqvc-fns	VSYGSLFDAT ma i dths-tg

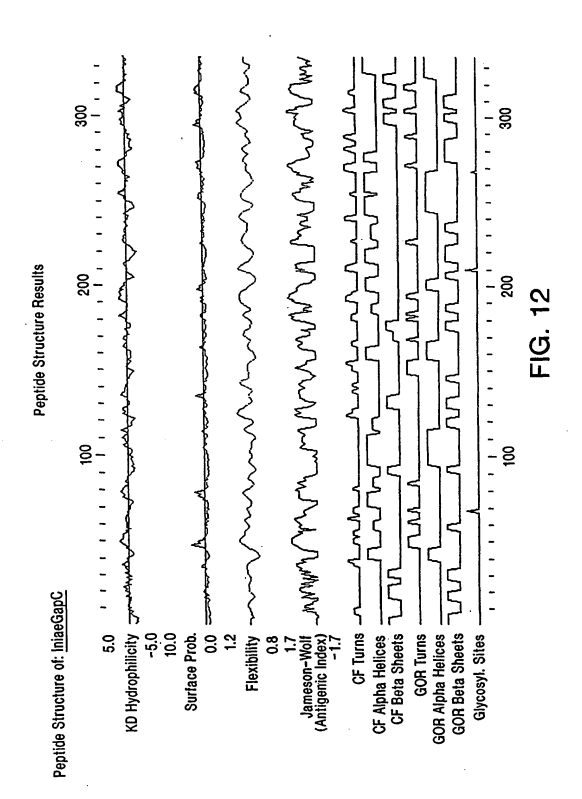
FIG. 7B

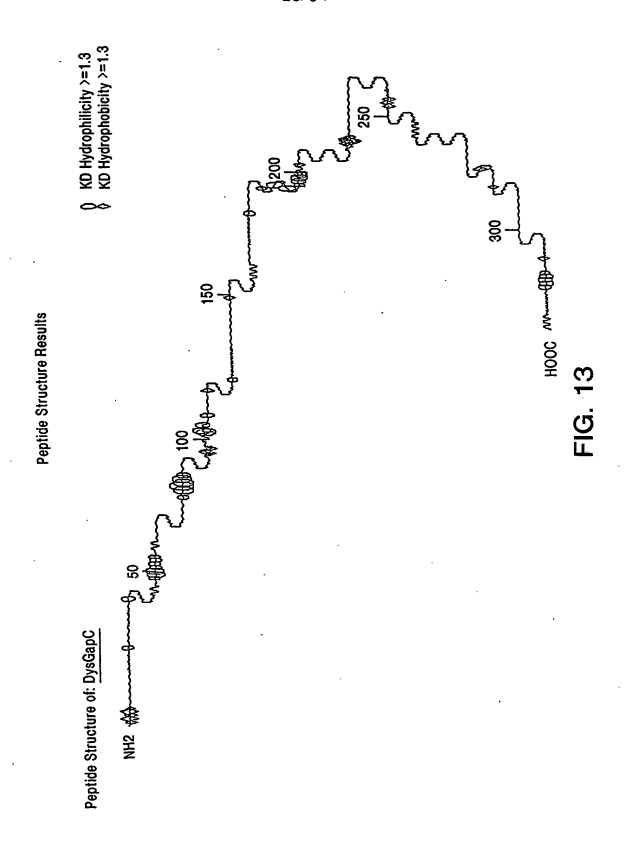


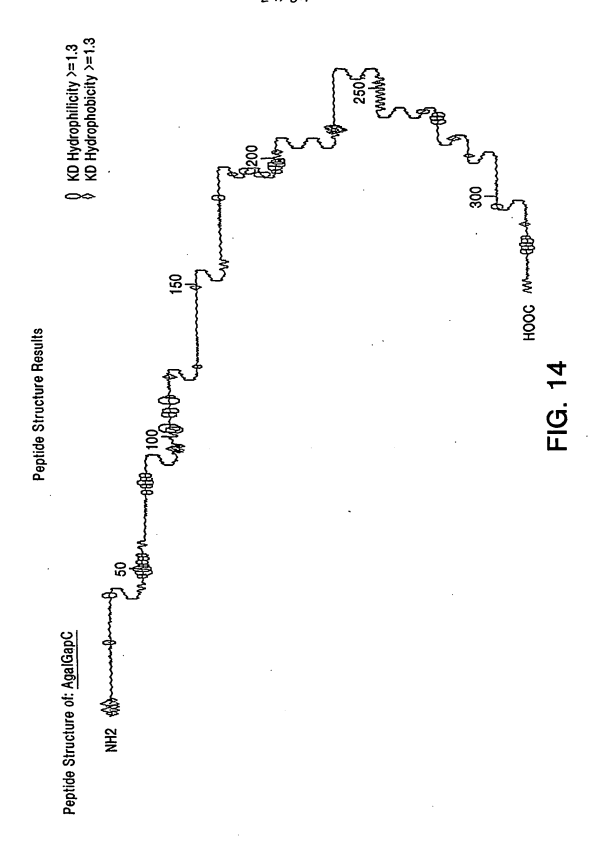


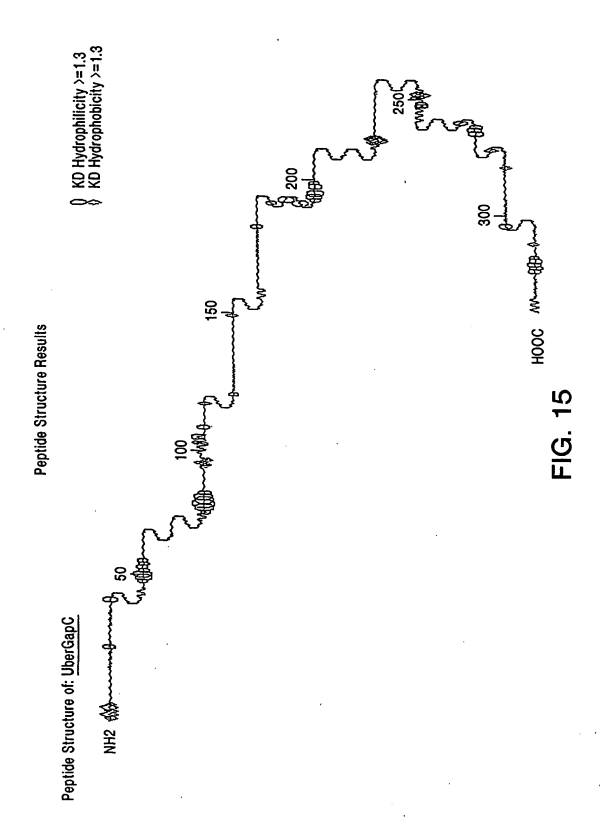


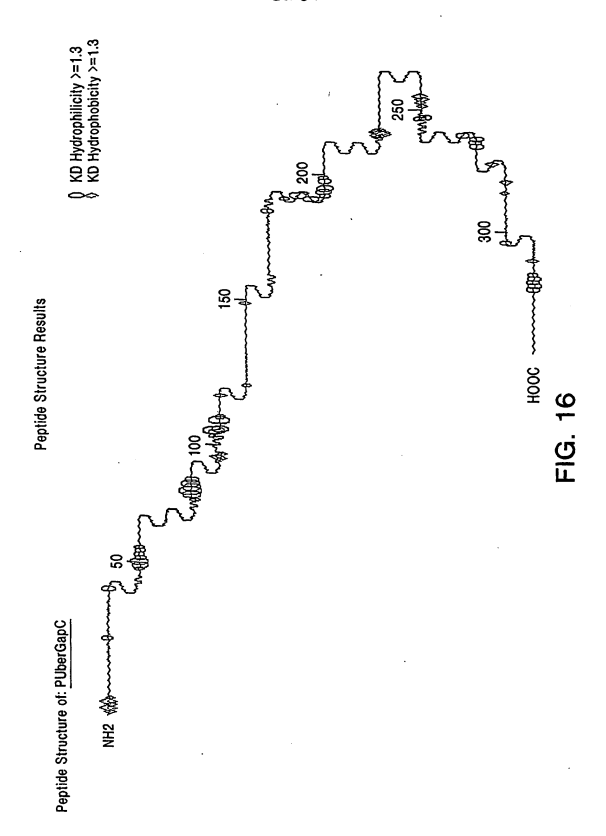


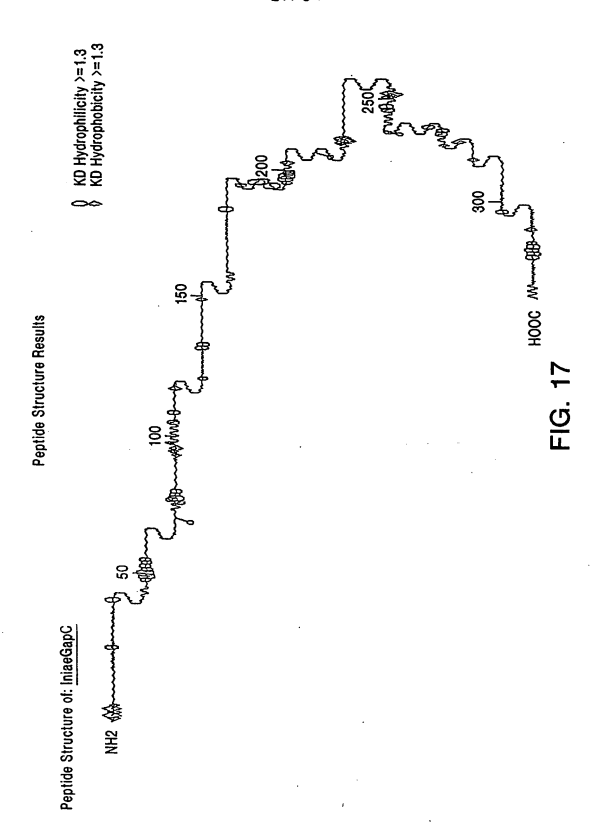












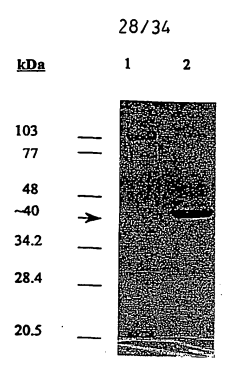


FIG. 18

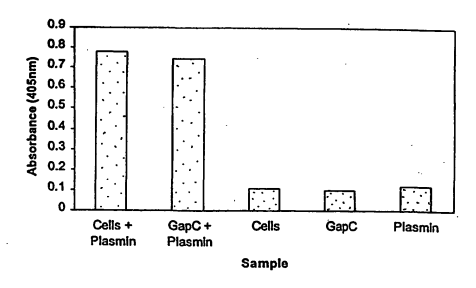


FIG. 19

29/34

Percentage of quaters infected with S. dysgalactiae per group

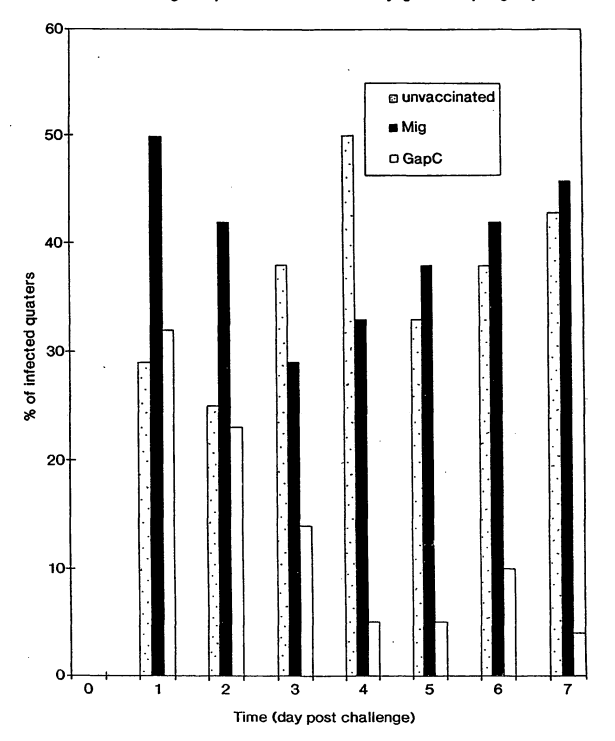


FIG. 20

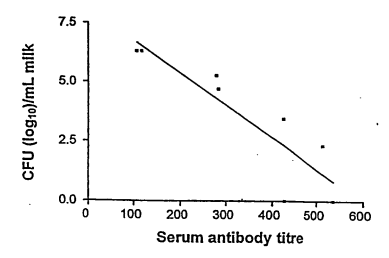


FIG. 21

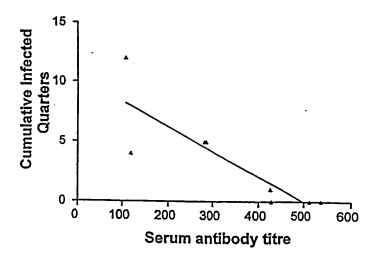


FIG. 22

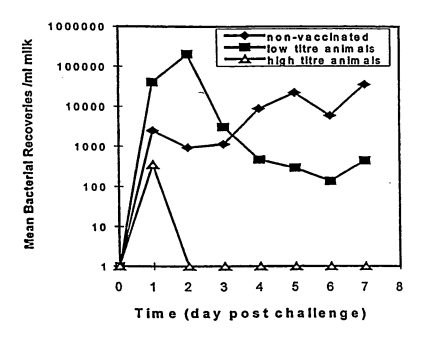


FIG. 23

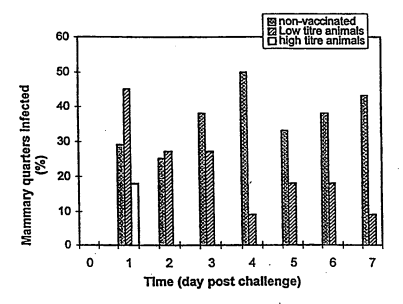


FIG. 24

Mean Somatic Cell Counts (SCC) per group

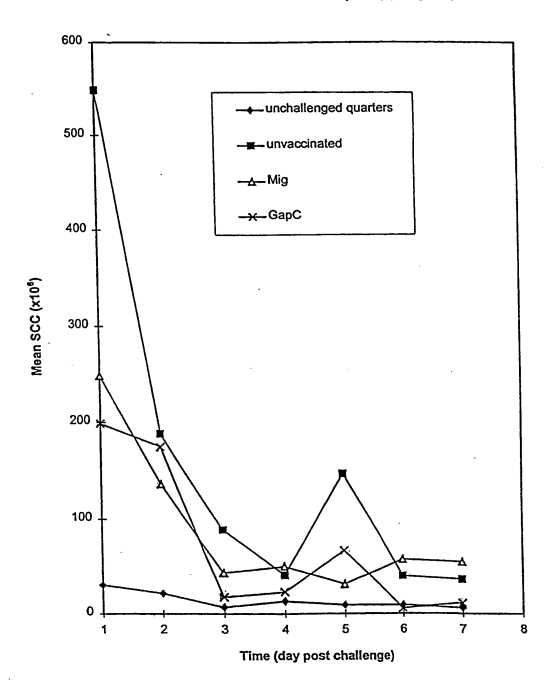


FIG. 25

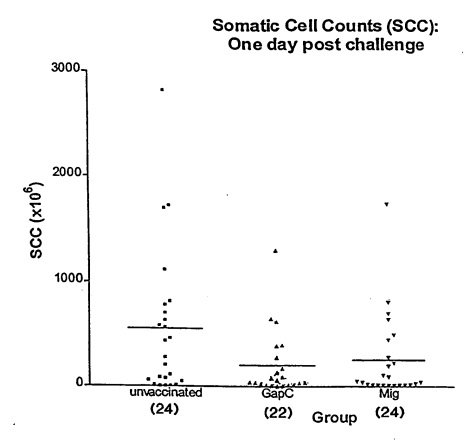


FIG. 26

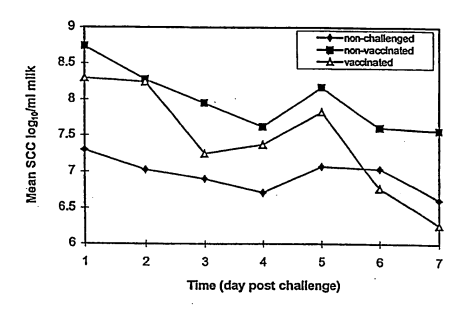


FIG. 27

SEQUENCE LISTING

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      Bolton, Alexandra J.
      Perez-Casal, Jose
      Fontaine, Michael
      Potter, Andrew A.
<120> IMMUNIZATION OF DAIRY CATTLE WITH GapC PROTEIN AGAINST
      STREPTOCOCCUS INFECTION
<130> 08-891814WO
<140>
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<160> 12
<170> PatentIn Ver. 2.0
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gca Ala	ttc Phe	cgt Arg	cgt Arg 20	att Ile	caa Gln	aat Asn	gtt Val	gaa Glu 25	ggt Gly	gtt Val	gaa Glu	gta Val	act Thr 30	cgt Arg	atc Ile	96
aac Asn	gac Asp	ctt Leu 35	aca Thr	gat Asp	cca Pro	aac Asn	atg Met 40	ctt Leu	gca Ala	cac His	ttg Leu	ttg Leu 45	aaa Lys	tac Tyr	gat Asp	144
Thr	Thr 50	Gln	gga Gly	Arg	Phe	Asp 55	Gly	Thr	Val	Glu	Val 60	Lys	Glu	Gly	Gly	192
Phe 65	Glu	Val	aac Asn	Gly	Asn 70	Phe	lle	Lys	Val	Ser 75	Ala	Glu	Arg	Asp	Pro 80	240
Glu	Asn	Ile	gac Asp	Trp 85	Ala	Thr	Asp	Gly	Val 90	Glu	Ile	Val	Leu	Glu 95	Ala	288
Thr	Gly	Phe	ttt Phe 100	Ala	Lys	Lys	Glu	Ala 105	Ala	Glu	Lys	His	Leu 110	His	Ala	336
Asn	Gly	Ala 115	aaa Lys	Lys	Val	Val	Ile 120	Thr	Ala	Pro	Gly	Gly 125	Asn	Asp	Val	384
			gtt Val													432
Thr 145	Val	Ile	tca Ser	Gly	Ala 150	Ser	Cys	Thr	Thr	Asn 155	Cys	Leu	Āla	Pro	Met 160	480
			ctt Leu													528
aca Thr			gct Ala 180													576
. –			gac Asp		_	-	_	_	_		_	-			-	624
			act Thr													672

210 215

ttg aat ggt aaa ctt gat ggt gct gca caa cgt gtt cct gtt cca act Leu Asn Gly Lys Leu Asp Gly Ala Ala Gln Arg Val Pro Val Pro Thr 225 235 gga tca gta act gag ttg gtt gta act ctt gat aaa aac gtt tct gtt 768 . Gly Ser Val Thr Glu Leu Val Val Thr Leu Asp Lys Asn Val Ser Val 245 gac gaa atc aac gct gct atg aaa gct gct tca aac gac agt ttc ggt Asp Glu Ile Asn Ala Ala Met Lys Ala Ala Ser Asn Asp Ser Phe Gly tac act gaa gat cca att gtt tct tca gat atc gta ggc gtg tca tac Tyr Thr Glu Asp Pro Ile Val Ser Ser Asp Ile Val Gly Val Ser Tyr 280 ggt tca ttg ttt gac gca act caa act aaa gtt atg gaa gtt gac gga 912 Gly Ser Leu Phe Asp Ala Thr Gln Thr Lys Val Met Glu Val Asp Gly 295 tca caa ttg gtt aaa gtt gta tca tgg tat gac aat gaa atg tct tac 960 Ser Gln Leu Val Lys Val Val Ser Trp Tyr Asp Asn Glu Met Ser Tyr 310 315 act gct caa ctt gtt cgt aca ctt gag tac ttt gcà aaa atc gct aaa 1008

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taa > 1011

330

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<211> 336

<212> PRT

<213> Streptococcus dysgalactiae

325

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Asn Asp Leu Thr Asp Pro Asn Met Leu Ala His Leu Leu Lys Tyr Asp 35 40 45

Thr Thr Gln Gly Arg Phe Asp Gly Thr Val Glu Val Lys Glu Gly Gly 50 55 60

Phe Glu Val Asn Gly Asn Phe Ile Lys Val Ser Ala Glu Arg Asp Pro 65 70 75 80

Glu Asn Ile Asp Trp Ala Thr Asp Gly Val Glu Ile Val Leu Glu Ala

90

95

Thr Gly Phe Phe Ala Lys Lys Glu Ala Ala Glu Lys His Leu His Ala 100 105 110

Asn Gly Ala Lys Lys Val Val Ile Thr Ala Pro Gly Gly Asn Asp Val 115 120 125

Lys Thr Val Val Phe Asn Thr Asn His Asp Ile Leu Asp Gly Thr Glu 130 135 140

Thr Val Ile Ser Gly Ala Ser Cys Thr Thr Asn Cys Leu Ala Pro Met 145 150 155 160

Ala Lys Ala Leu His Asp Ala Phe Gly Ile Gln Lys Gly Leu Met Thr
165 170 175

Thr Ile His Ala Tyr Thr Gly Asp Gln Met Ile Leu Asp Gly Pro His 180 185 190

Arg Gly Gly Asp Leu Arg Arg Ala Arg Ala Gly Ala Ala Asn Ile Val 195 200 205

Pro Asn Ser Thr Gly Ala Ala Lys Ala Ile Gly Leu Val Ile Pro Glu 210 215 220

Leu Asn Gly Lys Leu Asp Gly Ala Ala Gln Arg Val Pro Val Pro Thr 225 230 235 240

Gly Ser Val Thr Glu Leu Val Val Thr Leu Asp Lys Asn Val Ser Val 245 250 255

Asp Glu Ile Asn Ala Ala Met Lys Ala Ala Ser Asn Asp Ser Phe Gly
260 265 270

Tyr Thr Glu Asp Pro Ile Val Ser Ser Asp Ile Val Gly Val Ser Tyr
275 280 285

Gly Ser Leu Phe Asp Ala Thr Gln Thr Lys Val Met Glu Val Asp Gly 290 295 300

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Thr Ala Gln Leu Val Arg Thr Leu Glu Tyr Phe Ala Lys Ile Ala Lys 325 330 335

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5 5 59 Cgt Ctt 48
10 The second of
gca ttc cgt cgc atc caa aac gta gaa ggt gtt gaa gtt act cgt atc '96
Ala Phe Arg Arg Ile Gln Asn Val Glu Gly Val Glu Val Thr Arg Ile 20 25
20 Ash Val Glu Gly Val Glu Val Thr Arg 11
30
aac gac ctt aca gat cca aac atg ctt gca cac ttg ttg aaa tat gac 144
Asp Leu Thr Asp Pro Asp Met Lou 22
Asn Asp Leu Thr Asp Pro Asn Met Leu Ala His Leu Leu Lys Tyr Asp 40
aca agt
aca act caa ggt cgt ttc gac ggt act gtt gaa gtt aaa gaa ggt gga 192
Thr Thr Gln Gly Arg Phe Asp Gly Thr Val Glu Val Lys Glu Gly Gly 50 60
55 Thr Val Glu Val Lys Glu Gly Gly
tto gas one
ttc gaa gtt aac ggt caa ttt gtt aaa gtt tct gct gaa cgc gaa cca 240
Phe Glu Val Asn Gly Gln Phe Val Lys Val Ser Ala Glu Arg Glu Pro
70 Val Lys Val Ser Ala Glu Arg Glu Pro
75 75 80
gca aac att gac tgg gct act gat ggc gta gaa atc gtt ctt gaa gca 288
Ala Asn Ile Asp Trp Ala Thr Asp Gly Val Glu Ile Val Leu Glu Ala 85 90
85 2 2 4 Val Glu Ile Val Leu Glu Ala
act ggt ttg tt.
Thr Gly Phe Phe Ala Ser Lys Glu Lys Ala Gly Gln His Ile His Glu 100 105
100 The File Ala Ser Lys Glu Lys Ala Gly Gla Res - 336
105 This lie His Glu
aat got got and
Asn Gly Ala Lys Lys Val Val Ile Thr Ala Pro Gly Gly Asn Asp Val 115 120
115 bys val Val Ile Thr Ala Pro Gly Clar 384
120 THE ASP Val
ada ada att att
Lys Thr Val Val Phe Asn Thr Asn His Asp Ile Leu Asp Gly Thr Glu 130 135
130 Ash Thr Ash His Asp Ile Leu Asp Cly My 432
135 140
aca gtt atc tca ggt gct tca tgt act aca aac tgt ctt gct cca atg 480
Thr Val Ile Ser Gly Ala con generated aca aac tgt ctt gct cga at
Thr Val Ile Ser Gly Ala Ser Cys Thr Thr Asn Cys Leu Ala Pro Met
155 160
gct aaa gct tta caa gac aac ttt ggt gtt aaa caa ggt ttg atg act 528
Ala Lys Ala Leu Gln Asp Asn Phe Gly Val Lys Gln Gly Leu Met Thr
165 Ash Phe Gly Val Lys Gln Gly Leu Met Thr
170
,
act sta
act atc cac gca tac act ggt gac caa atg atc ctt gac gga cca cac 576
The His Ala Tyr Thr Gly Asp Gln Met Ile Lev Assault State Cac 576
Thr Ile His Ala Tyr Thr Gly Asp Gln Met Ile Leu Asp Gly Pro His
Cat gat and
cgt ggt ggt gac ctt cgt cgt gct cgt gca ggt gct gca aac atc gtt 624
195 Leu Arg Arg Ala Arg Ala Gly Ala Ala atc gtt 624
200 And Ash Ile Val
cct aac tca aat
cct aac tca act ggt gct gca aaa gct atc gga ctt gtt atc cca gaa 672
gga ctt gtt atc cca gaa 672
• • • • • • • • • • • • • • • • • • • •



Pro	Asn 210	Ser	Thr	Gly	Ala	Ala 215	Lys	Ala	Ile	Gly	Leu 220	Val	Ile	Pro	Glu	
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gga Gly	tca Ser	gta Val	act Thr	gaa Glu 245	ttg Leu	gtt Val	gca Ala	act Thr	ctt Leu 250	gaa Glu	aaa Lys	gac Asp	gta Val	act Thr 255	gtc Val	768
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ggt Gly	tca Ser 290	ttg Leu	ttt Phe	gat Asp	gct Ala	act Thr 295	caa Gln	act Thr	aaa Lys	Val	caa Gln 300	act Thr	gtt Val	gac Asp	ggt Gly	912
aac Asn 305	caa Gln	ttg Leu	gtt Val	aaa Lys	gtt Val 310	gtt Val	tca Ser	tgg Trp	tac Tyr	gat Asp 315	aac Asn	gaa Glu	atg Met	tca Ser	tac Tyr 320	960
act Thr	tca Ser	caa Gln	ctt Leu	gtt Val 325	cgt Arg	aca Thr	ctt Leu	Glu	tac Tyr 330	ttt Phe	gca Ala	aaa Lys	atc Ile	gct Ala 335	aaa Lys	1008
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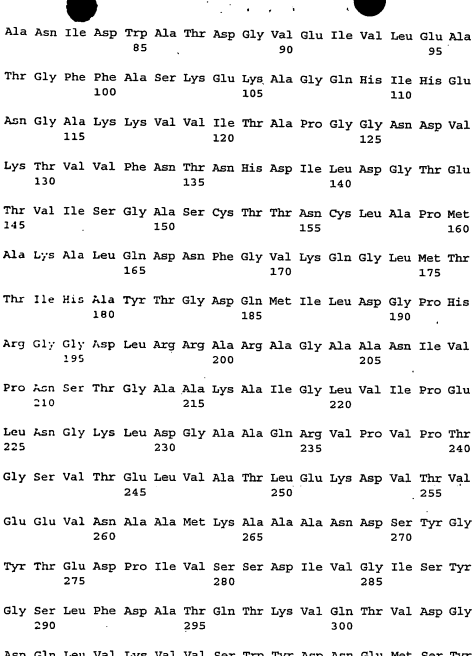
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Ala Phe Arg Arg Ile Gln Asn Val Glu Gly Val Glu Val Thr Arg Ile 20 25 30

Asn Asp Leu Thr Asp Pro Asn Met Leu Ala His Leu Leu Lys Tyr Asp 35. 40 45

Thr Thr Gln Gly Arg Phe Asp Gly Thr Val Glu Val Lys Glu Gly Gly 50 55 60

Phe Glu Val Asn Gly Gln Phe Val Lys Val Ser Ala Glu Arg Glu Pro 65 70 75 . 80



Asn Gln Leu Val Lys Val Val Ser Trp Tyr Asp Asn Glu Met Ser Tyr 305 310 315

Thr Ser Gln Leu Val Arg Thr Leu Glu Tyr Phe Ala Lys Ile Ala Lys

<210> 7

<211> 1011

<212> DNA

<213> Streptococcus uberis

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<221> CDS



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Ala	ttc Phe	Arg	Arg 20	Ile	Gln	Asn	Val	Glu 25	Gly	Val	Glu	Val	Thr 30	Arg	Ile	96
Asn	gat Asp	Leu 35	Thr	Asp	Pro	Asn	Met 40	Leu	Ala	His	Leu	Leu 45	Lys	Tyr	Asp	144
aca Thr	act Thr 50	caa Gln	ggt Gly	cgt Arg	ttc Phe	gac Asp 55	ggt Gly	aca Thr	gtt Val	gaa Glu	gtt Val 60	aaa Lys	gat Asp	ggt Gly	gga Gly	192
	gaa Glu															240
	aac Asn															288
	ggt Gly															336
	ggt															384
	act Thr 130		_						_			_			_	432
	gta Val				_		_				_		_		_	480
_	aaa Lys	_	_		_				_				_	_		528
	atc Ile															576
	ggt Gly	_			_	_		_			_	_			-	624

3

•	

									•							
cct Pro	aac Asn 210	tca Ser	act Thr	ggt Gly	gct Ala	gct Ala 215	aaa Lys	gca Ala	atc Ile	ggt Gly	ctt Leu 220	gta Val	atc Ile	cca Pro	gaa Glu	672
tta Leu 225	aat Asn	ggt Gly	aaa Lys	ctt Leu	gac Asp 230	ggt Gly	gct Ala	gca Ala	caa Gln	cgt Arg 235	gtt Val	cct Pro	gtt Val	cca Pro	act Thr 240	720
gga Gly	tca Ser	gta Val	act Thr	gaa Glu 245	tta Leu	gta Val	gca Ala	gtt Val	ctt Leu 250	gaa Glu	aaa Lys	gaa Glu	act Thr	tca Ser 255	gtt Val	768
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aat Asn 305	caa Gln	tta Leu	gtt Val	aaa Lys	gtt Val 310	gtt Val	tca Ser	tgg Trp	tat Tyr	gac Asp 315	aac Asn	gaa Glu	atg Met	tct Ser	tac Tyr 320	960
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taa									٠							1011

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<213> Streptococcus uberis

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Asn Asp Leu Thr Asp Pro Asn Met Leu Ala His Leu Leu Lys Tyr Asp 35 40 45

Thr Thr Gln Gly Arg Phe Asp Gly Thr Val Glu Val Lys Asp Gly Gly 50 55 60

Phe Glu Val Asn Gly Asn Phe Ile Lys Val Ser Ala Glu Lys Asp Pro 65 70 75 80



Glu Asn Ile Asp Trp Ala Thr Asp Gly Val Glu Ile Val Leu Glu Ala 85 Thr Gly Phe Phe Ala Lys Lys Ala Ala Ala Glu Lys His Leu His Ala 105 Asn Gly Ala Lys Lys Val Val Ile Thr Ala Pro Gly Gly Asp Asp Val . 120 Lys Thr Val Val Phe Asn Thr Asn His Asp Ile Leu Asp Gly Thr Glu 135 Thr Val Ile Ser Gly Ala Ser Cys Thr Thr Asn Cys Leu Ala Pro Met 150 155 Ala Lys Ala Leu Gln Asp Asn Phe Gly Val Lys Gln Gly Leu Met Thr 170 Thr Ile His Ala Tyr Thr Gly Asp Gln Met Ile Leu Asp Gly Pro His 185 Arg Gly Gly Asp Leu Arg Arg Ala Arg Ala Gly Ala Ser Asn Ile Val Pro Asn Ser Thr Gly Ala Ala Lys Ala Ile Gly Leu Val Ile Pro Glu Leu Asn Gly Lys Leu Asp Gly Ala Ala Gln Arg Val Pro Val Pro Thr 230 235 Gly Ser Val Thr Glu Leu Val Ala Val Leu Glu Lys Glu Thr Ser Val Glu Glu Ile Asn Ala Ala Met Lys Ala Ala Ala Asn Asp Ser Tyr Gly 265 Tyr Thr Glu Asp Pro Ile Val Ser Ser Asp Ile Ile Gly Met Ala Tyr 275 Gly Ser Leu Phe Asp Ala Thr Gln Thr Lys Val Gln Thr Val Asp Gly Asn Gln Leu Val Lys Val Val Ser Trp Tyr Asp Asn Glu Met Ser Tyr 310 Thr Ala Gln Leu Val Arg Thr Leu Glu Tyr Phe Ala Lys Ile Ala Lys 325 330

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gct	ttc	cgt	cgt	att	caa	aat	gta	gaa	ggt	gtt	gaa	gtt	act	cgc	atc	96
Ala	Phe	Arg	Arg	Ile	${\tt Gln}$	Asn	Val	Glu	Gly	Val	Glu	Val	Thr	Arg	Ile	
			20					25					30			

aac	gac	ctt	aca	gat	cca	aat	atg	ctt	gca	cac	ttg	tta	aaa	tac	gat	144
Asn																
		35					40					45	•	-	-	

aca	act	caa	ggt	cgt	ttt	gac	ggt	act	gta	gaa	gtt	aaa	gat	ggt	gga	192
Thr																
	50					55	_				60	•	-	_	-	

																240
Phe	Asp	Val	Asn	Gly	Lys	Phe	Ile	Lys	Val	Ser	Ala	Glu	Lys	Asp	Pro	
65					70					75		•			80	

gaa Glu									
		85.			90			95	

act	ggt	ttc	ttt	gct	aaa	aaa	gca	gct	gct	gaa	aaa	cat	tta	cat	gaa	336
Thr	Gly	Phe	Phe	Ala	Lys	Lys	Ala	Ala	Ala	Glu	Lys	His	Leu	His	Glu	
			100					105					110			

a	at	aat	act	222	222	att	att	atc	act	act	act	aat	~~=	rat	asc.	ata	-384
		_															-20-4
P	lsn	GIY	Ala	Lys	Lys	Val	Val	Ile	Thr	Ala	Pro	Gly	Gly	Asp	Asp	Val	
		•	115					120					125				•

aaa	aca	gtt	gta	ttt	aac	act	aac	cat	gat	atc	ctt	gat	gga	act	gaa	432
Lys	Thr	Val	Val	Phe	Asn	Thr	Asn	His	Asp	Ile	Leu	Asp	Gly	Thr	Glu	
	130					135					140					

aca	gtt	att	tca	ggt	gct.	tca	tgt	act	aca	aac	tgt	tta	gct	cca	atg	480
Thr	Val	Ile	Ser	Gly	Ala	Ser	Cys	Thr	Thr	Asn	Cys	Leu	Ala	Pro	Met	•
145					150					155					160	

gct	222	act	tta	caa	gat	aac	+++	aac	at a	222	caa	aat	tta	ato	act	528
900	uuu	300	cua	حببد	gac	uuc		990	gua	auu	caa	330	cca	aug	acc	720
Ala	Lvs	Ala	Leu	Gln	Asp	Asn	Phe	Glv	Val	Lvs	Gln	Glv	Leu	Met	Thr	
	-2				E			1		2		1				
				165		•			170					175		

aca	atc	cac	gct	tac	act	ggt	gat	caa	atg	ctt	ctt	gat	gga	cct	cac	576
Thr	Ile	His	Ala	Tyr	Thr	Gly	Asp	Gln	Met	Leu	Leu	Asp	Gly	${\tt Pro}$	His	
			180					185					190			

cgt Arg									624
	195			200			205		

cct aac tca act ggt gct gct aaa gca atc ggt ctt gtt atc cct gaa 672

WO 01/96381



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ggt Gly	tca Ser	gta Val	aca Thr	gaa Glu 245	tta Leu	gta Va'l	gca Ala	gtt Val	ctt Leu 250	aat Asn	aaa Lys	gaa Glu	act Thr	tca Ser 255	gta Val	768
gaa Glu	gaa Glu	att Ile	aac Asn 260	tca Ser	gta Val	atg Met	aaa Lys	gct Ala 265	gca Ala	gct Ala	aat Asn	gat Asp	tca Ser 270	tat Tyr	ggt Gly	816
tac Tyr	act Thr	gaa Glu 275	gat Asp	cca Pro	atc Ile	gta Val	tca Ser 280	tct Ser	gat Asp	atc Ile	gtt Val	ggt Gly 285	atg Met	tct Ser	ttc Phe	864
ggt Gly	tca Ser 290	tta Leu	ttc Phe	gat Asp	gct Ala	act Thr 295	caa Gln	act Thr	aaa Lys	gta Val	caa Gln 300	act Thr	gtt Val	gat Asp	gga Gly	912
aat Asn 305	caa Gln	tta Leu	gtt Val	aaa Lys	gtt Val 310	gtt Val	tca Ser	tgg Trp	tat Tyr	gac Asp 315	aat Asn	gaa Glu	atg Met	tct Ser	tac Tyr 320	960
act Thr	gct Ala	caa Gln	ctt Leu	gat Asp 325	cgt Arg	aca Thr	ctt Leu	gag Glu	tac Tyr 330	ttt Phe	gca Ala	aaa Lys	atc Ile	gct Ala 335	aaa Lys	1008
taa			• •									-				1011

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Met Val Val Lys Val Gly Ile Asn Gly Phe Gly Arg Ile Gly Arg Leu

1 10 15

Ala Phe Arg Arg Ile Gln Asn Val Glu Gly Val Glu Val Thr Arg Ile
20 25 30

Asn Asp Leu Thr Asp Pro Asn Met Leu Ala His Leu Leu Lys Tyr Asp 35 40 45

Thr Thr Gln Gly Arg Phe Asp Gly Thr Val Glu Val Lys Asp Gly Gly 50 55 60

Phe Asp Val Asn Gly Lys Phe Ile Lys Val Ser Ala Glu Lys Asp Pro 65 70 75 80



Glu Gln Ile Asp Trp Ala Thr Asp Gly Val Glu Ile Val Leu Glu Ala
85 90 95

Thr Gly Phe Phe Ala Lys Lys Ala Ala Ala Glu Lys His Leu His Glu 100 105 110

Asn Gly Ala Lys Lys Val Val Ile Thr Ala Pro Gly Gly Asp Asp Val 115 120 125

Lys Thr Val Val Phe Asn Thr Asn His Asp Ile Leu Asp Gly Thr Glu 130 135 140

Thr Val Ile Ser Gly Ala Ser Cys Thr Thr Asn Cys Leu Ala Pro Met 145 150 155 160

Ala Lys Ala Leu Gln Asp Asn Phe Gly Val Lys Gln Gly Leu Met Thr 165 170 175

Thr Ile His Ala Tyr Thr Gly Asp Gln Met Leu Leu Asp Gly Pro His 180 185 190

Arg Gly Gly Asp Leu Arg Arg Ala Arg Ala Gly Ala Asn Asn Ile Val 195 200 205

Pro Asn Ser Thr Gly Ala Ala Lys Ala Ile Gly Leu Val Ile Pro Glu 210 215 220

Leu Asn Gly Lys Leu Asp Gly Ala Ala Gln Arg Val Pro Val Pro Thr 225 230 235 240

Gly Ser Val Thr Glu Leu Val Ala Val Leu Asn Lys Glu Thr Ser Val 245 250 255

Glu Glu İle Asn Ser Val Met Lys Ala Ala Ala Asn Asp Ser Tyr Gly
260 265 270

Tyr Thr Glu Asp Pro Ile Val Ser Ser Asp Ile Val Gly Met Ser Phe 275 280 285

Gly Ser Leu Phe Asp Ala Thr Gln Thr Lys Val Gln Thr Val Asp Gly 290 295 300

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(19) World Intellectual Property Organization International Bureau





(43) International Publication Date 20 December 2001 (20.12.2001)

(10) International Publication Number WO 01/096381 A3

(51) International Patent Classification7: C07K 14/315, C12N 15/31, 15/70, 1/21, A61K 39/09, C07K 16/12, AGIP 314H. AGIK 31/7088, G01N 33/569, 33/577, 33/68

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- (21) International Application Number: PCT/CA01/00838
- (74) Agents: ERRATT, Judy, A. et al.; Gowlings Lafleur Henderson LLP, Suite 2600, 160 Elgin Street, Ottawa, Ontario K1P 1C3 (CA).

(84) Designated States (regional): European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC,

- (22) International Filing Date: 11 June 2001 (11.06.2001)
- (81) Designated States (national): CA, CO.

(25) Filing Language:

English

(26) Publication Language:

S7N 5L3 (CA)

English

(30) Priority Data:

66V211 022

12 June 2000 (12.06.2000) US

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- with international search report

NL, PT, SE, TR).

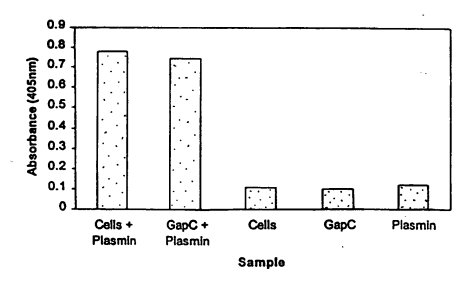
Published:

- (72) Inventors: BOLTON, Alexandra, J.; 115-1540
- (88) Date of publication of the international search report: 30 May 2003

29th Street N.W., Calgary, Alberta T2N 4M1 (CA), PEREZ-CASAL, Jose: 534 Avenue W.S., Saskatoon. Saskatchewan S7M 3G8 (CA). FONTAINE,

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(54) Title: IMMUNIZATION OF DAIRY CATTLE WITH GapC PROTEIN AGAINST STREPTOCOCCUS INFECTION



01/096381 A3 (57) Abstract: The GapC plasmin binding protein genes of Streptococcus dysgalactiae (S. dysgalactiae), Streptococcus agalactiae (S. agalactiae). Streptococcus uberis (S. uberis), Streptococcus parauberis (S. parauberis), and Streptococcus iniae (S.iniae) are described, as well as the recombinant production of the GapC proteins therefrom. Also described is the use of the GapC proteins from those species in vaccine compositions to prevent or treat bacterial infections in general, and mastitis in particular.

INTERNATIONAL SEARCH REPORT

al Application No r/CA 01/00838

A. CLASSIFICATION OF SUITE TO TO CO7K14/315

C12N15/31 C07K16/12 A61P31/04 G01N33/68

C12N15/70 C12N1/21 A61K31/7088 G01N33/569 A61K39/09 G01N33/577

According to International Patent Classification (IPC) or to both national classification and IPC

Minimum documentation searched (classification system followed by classification symbols) IPC 7 C07K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

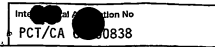
Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

BIOSIS, EMBASE, WPI Data, PAJ, EPO-Internal, SEQUENCE SEARCH

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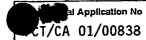
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